

## 2024 Prior Authorization Medical Necessity Guidelines

Effective: July 1, 2024 Updated: July 1, 2024

These guidelines were updated on July 1, 2024. For more recent information or other questions, please contact Mass General Brigham Health Plan Customer Service team. Visit **MassGeneralBrighamAdvantage.org/Rx-information** for the most up-to-date information on Medicare Part D drug coverage.

You can reach our Customer Service team by calling: **855-833-3668** (TTY: 711)

October 1 – March 31 8:00 AM to 8:00 PM EST, Monday through Sunday

April 1 – September 30 8:00 AM to 8:00 PM EST, Monday through Friday

Mass General Brigham Advantage Secure (HMO-POS) Mass General Brigham Advantage (PPO), and Mass General Brigham Advantage Premier (PPO)

Mass General Brigham Health Plan is a Medicare Advantage organization with a Medicare contract offering HMO-POS and PPO plans. Enrollment in Mass General Brigham Health Plan depends on contract renewal.

PA Criteria

**Prior Authorization Group ABIRATERONE** 

**Drug Names** ABIRATERONE ACETATE

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Node-positive (N1), non-metastatic (M0) prostate cancer and very-high-risk prostate

cancer.

**Exclusion Criteria** 

**Required Medical Information** 

The requested drug will be used in combination with a gonadotropin-releasing hormone

(GnRH) analog or after bilateral orchiectomy.

Age Restrictions

**Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria

**ADEMPAS Prior Authorization Group ADEMPAS Drug Names** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** 

For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group 1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood units. For chronic thromboembolic pulmonary hypertension (CTEPH) (WHO Group 4): 1) Patient has persistent or recurrent CTEPH after pulmonary endarterectomy (PEA), OR 2) Patient has inoperable CTEPH with the diagnosis confirmed by right heart catheterization AND by computed tomography (CT), magnetic resonance imaging (MRI), or pulmonary angiography.

Age Restrictions **Prescriber Restrictions** 

**Coverage Duration** Plan Year

Other Criteria

Updated 07/01/2024 1 Prior Authorization GroupADLARITYDrug NamesADLARITY

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Vascular dementia

Exclusion Criteria -

**Required Medical Information** Patient is unable to take oral dosage forms (e.g., difficulty swallowing tablets or

capsules). For dementia of the Alzheimer's type: the patient has experienced an inadequate response, intolerance, or the patient has a contraindication to rivastigmine

transdermal patch.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupAEMCOLODrug NamesAEMCOLO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of Traveler's diarrhea.

Age Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group AGAMREE
Drug Names AGAMREE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the treatment of Duchenne muscular dystrophy (DMD): 1) The diagnosis was

confirmed by genetic testing identifying a disease-causing mutation of the DMD gene AND 2) The patient has tried prednisone and experienced any of the following a)

unmanageable and/or clinically significant weight gain/obesity, b) unmanageable and/or

clinically significant psychiatric/behavioral issues such as abnormal behavior,

aggression, or irritability, or c) clinically significant growth stunting.

**Age Restrictions** 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupAIMOVIGDrug NamesAIMOVIG

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the preventive treatment of migraine, initial: 1) The patient experienced an

inadequate treatment response with a 4-week trial of any one of the following: antiepileptic drugs (AEDs), beta-adrenergic blocking agents, antidepressants OR 2) The patient experienced an intolerance or has a contraindication that would prohibit a 4-week trial of any one of the following: antiepileptic drugs (AEDs), beta-adrenergic blocking agents, antidepressants. For preventive treatment of migraine, continuation: The patient received at least 3 months of treatment with the requested drug, and the

patient had a reduction in migraine days per month from baseline.

Age Restrictions Prescriber Restrictions -

**Coverage Duration** Initial: 3 months, Continuation: Plan Year

Other Criteria -

**Prior Authorization Group** AKEEGA **Drug Names** AKEEGA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** The requested drug will be used in combination with a gonadotropin-releasing hormone

(GnRH) analog or after bilateral orchiectomy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ALECENSA
Drug Names ALECENSA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent ALK-positive non-small cell lung cancer (NSCLC), brain metastases from

ALK-positive NSCLC, ALK-positive anaplastic large-cell lymphoma.

Exclusion Criteria -

**Required Medical Information** For non-small cell lung cancer (NSCLC): the disease is recurrent, advanced, or

metastatic.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group

ALPHA1-PROTEINASE INHIBITOR

**Drug Names** ARALAST NP, PROLASTIN-C, ZEMAIRA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For alpha1-proteinase inhibitor deficiency: Patient must have 1) clinically evident

emphysema, AND 2) pretreatment serum alpha1-proteinase inhibitor level less than 11

micromol/L (80 mg/dL by radial immunodiffusion or 50 mg/dL by nephelometry).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ALUNBRIG
Drug Names ALUNBRIG

**PA Indication Indicator**All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent anaplastic lymphoma kinase (ALK)-positive non-small cell lung cancer

(NSCLC), brain metastases from ALK-positive NSCLC, inflammatory myofibroblastic

tumors (IMT) with ALK translocation.

Exclusion Criteria -

**Required Medical Information** For non-small cell lung cancer (NSCLC): 1) the disease is recurrent, advanced, or

metastatic AND 2) the disease is anaplastic lymphoma kinase (ALK)-positive.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group ALVAIZ ALVAIZ Drug Names** 

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** 

For chronic or persistent immune thrombocytopenia (ITP) (new starts): 1) Patient (pt) has experienced an inadequate treatment response or is intolerant to a prior therapy such as corticosteroids or immunoglobulins, AND 2) Untransfused platelet (plt) count at any point prior to the initiation of the requested medication is less than 30.000/mcL OR 30,000-50,000/mcL with symptomatic bleeding or risk factor(s) for bleeding (e.g., undergoing a medical or dental procedure where blood loss is anticipated. comorbidities such as peptic ulcer disease and hypertension, anticoagulation therapy, profession or lifestyle that predisposes pt to trauma). For ITP (continuation): plt count response to the requested drug: 1) Current plt count is less than or equal to 200,000/mcL, OR 2) Current plt count is greater than 200,000/mcL to less than or equal to 400,000/mcL and dosing will be adjusted to a plt count sufficient to avoid clinically important bleeding. For thrombocytopenia associated with chronic hepatitis C (new starts): the requested drug is used for initiation and maintenance of interferon-based therapy. For thrombocytopenia associated with chronic hepatitis C (continuation): pt is receiving interferon-based therapy. For severe aplastic anemia (AA) (new starts): Pt had an insufficient response to immunosuppressive therapy.

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Other Criteria

HCV: 6mo, ITP/AA initial: 6mo, ITP reauth: Plan Year, AA reauth: APR-Plan Year, IPR-16 wks

For severe AA (continuation): 1) Current plt count is 50,000-200,000/mcL, OR 2) Current plt count is less than 50.000/mcL and pt has not received appropriately titrated therapy for at least 16 weeks, OR 3) Current plt count is less than 50,000/mcL and pt is transfusion-independent, OR 4) Current plt count is greater than 200,000/mcL to less than or equal to 400,000/mcL and dosing will be adjusted to achieve and maintain an appropriate target plt count. APR: adequate platelet response (greater than 50,000/mcL), IPR: inadequate platelet response (less than 50,000/mcL).

Updated 07/01/2024 5 Prior Authorization Group AMBRISENTAN Drug Names AMBRISENTAN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group

1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood

units.

Age Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ARCALYST Drug Names ARCALYST

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Prevention of gout flares in patients initiating or continuing urate-lowering therapy.

Exclusion Criteria -

**Required Medical Information** For prevention of gout flares in patients initiating or continuing urate-lowering therapy

(e.g., allopurinol) (new starts): 1) two or more gout flares within the previous 12 months, AND 2) inadequate response, intolerance, or contraindication to maximum tolerated doses of a non-steroidal anti-inflammatory drug (NSAID) and colchicine, AND 3) concurrent use with urate-lowering therapy. For prevention of gout flares in patients initiating or continuing urate-lowering therapy (e.g., allopurinol) (continuation): 1) patient must have achieved or maintained a clinical benefit (i.e., a fewer number of gout attacks or fewer flare days) compared to baseline, AND 2) continued use of

urate-lowering therapy concurrently with the requested drug. For recurrent pericarditis: patient must have had an inadequate response, intolerance, or contraindication to

maximum tolerated doses of an NSAID and colchicine.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupARIKAYCEDrug NamesARIKAYCE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupARMODAFINILDrug NamesARMODAFINIL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For excessive sleepiness associated with narcolepsy: The diagnosis has been

confirmed by sleep lab evaluation. For excessive sleepiness associated with

obstructive sleep apnea (OSA): The diagnosis has been confirmed by

polysomnography.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group AUGTYRO
Drug Names AUGTYRO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group AURYXIA
Drug Names AURYXIA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** The requested drug is not being prescribed for treatment of iron deficiency anemia in

adult patients with chronic kidney disease not on dialysis.

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** AUSTEDO

**Drug Names** AUSTEDO, AUSTEDO XR, AUSTEDO XR PATIENT TITRAT

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Tourette's syndrome

Exclusion Criteria -

Required Medical Information - Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group AYVAKIT
Drug Names AYVAKIT

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Myeloid and lymphoid neoplasms with eosinophilia, gastrointestinal stromal tumor (GIST) for unresectable, recurrent, or metastatic disease without platelet-derived

growth factor receptor alpha (PDGFRA) exon 18 mutation.

Exclusion Criteria
Required Medical Information

Off-label Uses

For myeloid and lymphoid neoplasms with eosinophilia, the patient meets all of the following criteria: 1) The disease is FIP1L1- PDGFRA rearrangement-positive, AND 2) The disease harbors a PDGFRA D842A mutation, AND 3) The disease is resistant to imatinib. For GIST, the patient meets either of the following criteria: 1) The disease harbors PDGFRA exon 18 mutation, including PDGFRA D842V mutations, OR 2) The requested drug will be used after failure on at least two Food and Drug Administration (FDA)-approved therapies in unresectable, recurrent, or metastatic disease without PDGFRA exon 18 mutation. For systemic mastocytosis: 1) The patient has a diagnosis of indolent systemic mastocytosis or advanced systemic mastocytosis (including aggressive systemic mastocytosis [ASM], systemic mastocytosis with associated hematological neoplasm [SM-AHN], and mast cell leukemia [MCL]) AND 2) The patient has a platelet count of greater than or equal to 50,000/microliter (mcL).

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group Drug Names** 

B VS. D

ACETYLCYSTEINE. ALBUTEROL SULFATE. AMINOSYN II. AMINOSYN-PF 7%. APREPITANT, ARFORMOTEROL TARTRATE, ASTAGRAF XL, AZATHIOPRINE, BENDEKA, BIVIGAM, BUDESONIDE, CARBOPLATIN, CISPLATIN, CLINIMIX 4.25%/DEXTROSE 1, CLINIMIX 4.25%/DEXTROSE 5, CLINIMIX 5%/DEXTROSE 15%, CLINIMIX 5%/DEXTROSE 20%, CLINISOL SF 15%, CROMOLYN SODIUM, CYCLOPHOSPHAMIDE, CYCLOPHOSPHAMIDE MONOHYDR, CYCLOSPORINE. CYCLOSPORINE MODIFIED. DOXORUBICIN HYDROCHLORIDE. DRONABINOL. EMEND, ENGERIX-B. ENVARSUS XR. EVEROLIMUS, FLEBOGAMMA DIF. FORMOTEROL FUMARATE, FREAMINE III, GAMASTAN, GAMMAGARD LIQUID, GAMMAGARD S/D IGA LESS TH. GAMMAKED. GAMMAPLEX. GAMUNEX-C. GENGRAF, GRANISETRON HYDROCHLORIDE, HEPLISAV-B, IMOVAX RABIES (H.D.C.V.), INTRALIPID, IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE/ALBUT, JYLAMVO, LEVALBUTEROL, LEVALBUTEROL HCL, LEVALBUTEROL HYDROCHLORID, LEVOCARNITINE, METHOTREXATE, METHOTREXATE SODIUM, MYCOPHENOLATE MOFETIL, MYCOPHENOLIC ACID DR, NUTRILIPID, OCTAGAM, ONDANSETRON HCL, ONDANSETRON HYDROCHLORIDE, ONDANSETRON ODT, OXALIPLATIN, PANZYGA, PARAPLATIN, PENTAMIDINE ISETHIONATE, PLENAMINE, PREHEVBRIO, PREMASOL, PRIVIGEN, PROGRAF, PROSOL, PULMOZYME, RABAVERT, RECOMBIVAX HB, SIROLIMUS, TACROLIMUS, TENIVAC, TOBRAMYCIN, TPN ELECTROLYTES, TRAVASOL, TREXALL, TROPHAMINE, VARUBI, XATMEP, YUPELRI, ZOLEDRONIC ACID

PA Indication Indicator

Off-label Uses **Exclusion Criteria Required Medical Information** 

Other Criteria

**Age Restrictions Prescriber Restrictions Coverage Duration** 

All Medically-accepted Indications

N/A

This drug may be covered under Medicare Part B or D depending upon the circumstances. Information may need to be submitted describing the use and setting of

the drug to make the determination.

Updated 07/01/2024 10 Prior Authorization GroupBAFIERTAMDrug NamesBAFIERTAM

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBALVERSADrug NamesBALVERSA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For urothelial carcinoma: 1) disease has susceptible fibroblast growth factor receptor 3

(FGFR3) or fibroblast growth factor receptor 2 (FGFR2) genetic alterations AND 2) the requested drug will be used as subsequent therapy for any of the following: a) locally advanced or metastatic urothelial carcinoma, b) recurrent primary carcinoma of the urethra, c) stage II-IV urothelial carcinoma of the bladder, d) urothelial carcinoma of the bladder with metastatic or local recurrence post cystectomy, or e) urothelial carcinoma of the bladder with muscle invasive local recurrence or persistent disease in a

or the biadder with muscle invasive local recurrence or persistent disease in

preserved bladder.

Age Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBANZELDrug NamesRUFINAMIDE

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria - Required Medical Information -

Age Restrictions 1 year of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBENLYSTADrug NamesBENLYSTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

Exclusion Criteria For pa

**Required Medical Information** 

For patients new to therapy: severe active central nervous system lupus.

For systemic lupus erythematosus (SLE): 1) patient is currently receiving a stable standard therapy regimen (e.g., corticosteroid, antimalarial, or NSAIDs) for SLE, OR 2) patient has experienced an intolerance or has a contraindication to standard therapy regimen for SLE. For lupus nephritis: 1) patient is currently receiving a stable standard therapy regimen (e.g., corticosteroid, cyclophosphamide, mycophenolate mofetil, or azathioprine) for lupus nephritis OR 2) patient has experienced an intolerance or has a contraindication to standard therapy regimen for lupus nephritis.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group BERINERT Drug Names BERINERT

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** 

For acute angioedema attacks due to hereditary angioedema (HAE): Patient meets either of the following: 1) the patient has HAE with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing OR 2) the patient has HAE with normal C1 inhibitor confirmed by laboratory testing and one of the following: a) the patient tested positive for an F12, angiopoietin-1, plasminogen, kininogen-1 (KNG1), heparan

sulfate-glucosamine 3-O-sulfotransferase 6 (HS3ST6), or myoferlin (MYOF) gene mutation OR b) the patient has a family history of angioedema and the angioedema was refractory to a trial of high-dose antihistamine therapy for at least one month.

Age Restrictions 5 years of age or older

**Prescriber Restrictions**Prescribed by or in consultation with an immunologist, allergist, or rheumatologist **Coverage Duration**Plan Year

Other Criteria -

Prior Authorization GroupBESREMIDrug NamesBESREMI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBETASERONDrug NamesBETASERON

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** BOSENTAN

Drug NamesBOSENTAN, TRACLEERPA Indication IndicatorAll FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group

1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood

units.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group BOSULIF
Drug Names BOSULIF

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Philadelphia chromosome positive B-cell acute lymphoblastic leukemia (Ph+ B-ALL),
myeloid and/or lymphoid neoplasms with eosinophilia and ABL1 rearrangement in the

chronic phase or blast phase

Exclusion Criteria
Required Medical Information

For chronic myeloid leukemia (CML), including patients newly diagnosed with CML and patients who have received a hematopoietic stem cell transplant: 1) Diagnosis was confirmed by detection of the Philadelphia chromosome or BCR-ABL gene, AND 2) If patient experienced resistance to an alternative tyrosine kinase inhibitor, patient is negative for all of the following mutations: T315I, G250E, V299L, and F317L, AND 3) patient has experienced resistance or intolerance to imatinib or dasatinib. For B-ALL including patient who have received hematopoietic stem cell transplant: 1) Diagnosis was confirmed by detection of the Philadelphia chromosome or BCR-ABL gene, and 2) If patient experienced resistance to an alternative tyrosine kinase inhibitor, patient is negative for all of the following mutations: T315I, G250E, V299L, and F317L.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBRAFTOVIDrug NamesBRAFTOVI

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Adjuvant systemic therapy for cutaneous melanoma, appendiceal adenocarcinoma

Exclusion Criteria -

For colorectal cancer (including appendiceal adenocarcinoma): 1) Tumor is positive for BRAF V600E mutation, AND 2) The requested drug will be used for either of the following: a) subsequent therapy for advanced or metastatic disease, b) primary treatment for unresectable metachronous metastases. For melanoma: 1) Tumor is positive for BRAF V600 activating mutation (e.g., V600E or V600K), AND 2) The requested drug will be used as a single agent or in combination with binimetinib, AND 3) The requested drug will be used for either of the following: a) unresectable, limited resectable, or metastatic disease, b) adjuvant systemic therapy.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBRIVIACTDrug NamesBRIVIACT

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For treatment of partial-onset seizures (i.e., focal-onset seizures): 1) The patient has

experienced an inadequate treatment response, intolerance, or has a contraindication to a generic anticonvulsant AND 2) the patient has experienced an inadequate

treatment response, intolerance, or has a contraindication to any of the following:

Aptiom (if 4 years of age or older), Xcopri (if 18 years of age or older), Spritam (if 4

years of age or older).

Age Restrictions 1 month of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupBRUKINSADrug NamesBRUKINSA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** BYDUREON

**Drug Names** BYDUREON BCISE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information -

**Age Restrictions** 10 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria The Prior Authorization only applies to patients whose claim is not submitted with an

ICD-10 code indicating a diagnosis of type 2 diabetes mellitus OR to patients who do not have a history of an antidiabetic drug (EXCLUDING glucagon-like peptide receptor agonists [GLP-1 RAs] and combination glucose-dependent insulinotropic polypeptide

[GIP] and GLP-1 RAs).

**Prior Authorization Group Drug Names**BYETTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria The Prior Authorization only applies to patients whose claim is not submitted with an

ICD-10 code indicating a diagnosis of type 2 diabetes mellitus OR to patients who do not have a history of an antidiabetic drug (EXCLUDING glucagon-like peptide receptor agonists [GLP-1 RAs] and combination glucose-dependent insulinotropic polypeptide

[GIP] and GLP-1 RAs).

**Prior Authorization Group** BYLVAY

**Drug Names**BYLVAY, BYLVAY (PELLETS) **PA Indication Indicator**All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For treatment of pruritis in progressive familial intrahepatic cholestasis (PFIC) (initial

requests): 1) diagnosis of PFIC has been confirmed by genetic testing, 2) the patient does not have PFIC type 2 with ABCB11 variants resulting in non-functional or

complete absence of bile salt export pump protein (BSEP-3), 3) the patient does not have any other concomitant liver disease, AND 4) the patient has not received a liver transplant. For treatment of pruritis in PFIC (continuation requests): the patient has experienced benefit from therapy (for example, improvement in pruritis). For treatment

of cholestatic pruritus with Alagille Syndrome (ALGS) (continuation): the patient has

experienced benefit from therapy (for example, improvement in pruritis). For PFIC: 3 months of age or older, For ALGS: 12 months of age or older Prescribed by or in consultation with a hepatologist or gastroenterologist

**Coverage Duration** Initial: 6 months, Continuation: Plan Year

Other Criteria -

Age Restrictions

**Prescriber Restrictions** 

Prior Authorization Group CABOMETYX
Drug Names CABOMETYX

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Non-small cell lung cancer, Ewing sarcoma, osteosarcoma, gastrointestinal stromal

tumor, endometrial carcinoma

Exclusion Criteria
Required Medical Information

For renal cell carcinoma: The disease is advanced, relapsed, or stage IV. For non-small cell lung cancer: 1) the disease is rearranged during transfection (RET) positive AND 2) the disease is recurrent, advanced, or metastatic. For hepatocellular carcinoma: the requested drug will be used as subsequent treatment. For gastrointestinal stromal tumor (GIST): The patient meets either of the following: 1) the disease is unresectable, recurrent/progressive, or metastatic AND the patient has failed a FDA-approved therapy (e.g., imatinib, sunitinib, regorafenib, ripretinib) OR 2) the requested drug will be used for palliation of symptoms if previously tolerated and effective. For Ewing sarcoma and osteosarcoma: the requested drug will be used as subsequent therapy. For differentiated thyroid cancer (DTC) (follicular, papillary, Hurthle cell): 1) The disease is locally advanced or metastatic disease, 2) the disease has progressed after a vascular endothelial growth factor receptor (VEGFR)- targeted therapy, AND 3) the patient is refractory to radioactive iodine therapy (RAI) or ineligible for RAI.For endometrial carcinoma: 1) the disease is recurrent or metastatic AND 2) the requested drug will be used as subsequent therapy.

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group CALQUENCE
Drug Names CALQUENCE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Waldenstrom macroglobulinemia (lymphoplasmacytic lymphoma), marginal zone lymphoma (including extranodal marginal zone lymphoma of the stomach, extranodal marginal zone lymphoma of nongastric sites, nodal marginal zone lymphoma, splenic

marginal zone lymphoma)

Exclusion Criteria

Off-label Uses

**Required Medical Information** For marginal zone lymphoma (including extranodal marginal zone lymphoma of the

stomach, extranodal marginal zone lymphoma of nongastric sites, nodal marginal zone lymphoma, and splenic marginal zone lymphoma): the requested drug is being used for

the treatment of relapsed, refractory, or progressive disease.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCAMZYOSDrug NamesCAMZYOS

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For obstructive hypertropic cardiomyopathy: 1) before initiating therapy, patient has left

ventricular ejection fraction (LVEF) of 55 percent or greater AND 2) patient has New

York Heart Association (NYHA) class II-III symptoms.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCAPRELSADrug NamesCAPRELSA

PA Indication IndicatorAll FDA-approved Indications, Some Medically-accepted IndicationsOff-label UsesDifferentiated thyroid carcinoma: papillary, follicular, and Hurthle cell.

Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** CARBAGLU

**Drug Names** CARGLUMIC ACID

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For N-acetylglutamate synthase (NAGS) deficiency: Diagnosis of NAGS deficiency was

confirmed by enzymatic, biochemical, or genetic testing.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCAYSTONDrug NamesCAYSTON

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For treatment of respiratory symptoms in cystic fibrosis patients: 1) Pseudomonas

aeruginosa is present in the patient's airway cultures. OR 2) The patient has a history

of pseudomonas aeruginosa infection or colonization in the airways.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group CERDELGA
Drug Names CERDELGA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For type 1 Gaucher disease (GD1): 1) Diagnosis was confirmed by an enzyme assay

demonstrating a deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing, and 2) Patient's CYP2D6 metabolizer status has been established using an FDA-cleared test, and 3) Patient is a CYP2D6 extensive metabolizer, an intermediate

metabolizer, or a poor metabolizer.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCEREZYMEDrug NamesCEREZYME

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Type 2 Gaucher disease, Type 3 Gaucher disease.

Exclusion Criteria -

**Required Medical Information** For Gaucher disease: Diagnosis was confirmed by an enzyme assay demonstrating a

deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCHOLBAMDrug NamesCHOLBAM

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information For bile acid synthesis disorders due to single enzyme defects (SEDs) and adjunctive

treatment of peroxisomal disorders (PDs): Diagnosis was confirmed by mass

spectrometry or other biochemical or genetic testing. For bile acid synthesis disorders due to SEDs and adjunctive treatment of PDs, continuation of therapy: Patient has

achieved and maintained improvement in liver function.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Initial: 6 months, Continuation: Plan Year

Other Criteria -

Prior Authorization Group CIBINQO
Drug Names CIBINQO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For atopic dermatitis (AD), continuation of therapy: Patient achieved or maintained

positive clinical response.

Age Restrictions 12 years of age or older

Prescriber Restrictions -

**Coverage Duration** Initial: 4 months, Continuation: Plan Year

Other Criteria -

Prior Authorization Group CINRYZE Drug Names CINRYZE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the prevention of acute angioedema attacks due to hereditary angioedema (HAE):

Patient meets either of the following: 1) the patient has hereditary angioedema (HAE) with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing OR 2) the patient has hereditary angioedema with normal C1 inhibitor confirmed by laboratory testing and either of the following: a) Patient tested positive for an F12, angiopoietin-1, plasminogen, kininogen-1 (KNG1), heparan sulfate-glucosamine 3-O-sulfotransferase 6 (HS3ST6), or myoferlin (MYOF) gene mutation OR b) Patient has a family history of angioedema and the angioedema was refractory to a trial of high-dose antihistamine

therapy for at least one month.

**Age Restrictions** 6 years of age or older

Prescriber Restrictions Prescribed by or in consultation with an immunologist, allergist, or rheumatologist

Coverage Duration Plan Year
Other Criteria -

Prior Authorization Group CLOBAZAM
Drug Names CLOBAZAM

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Seizures associated with Dravet syndrome

Exclusion Criteria - Required Medical Information -

Age Restrictions Seizures associated with Lennox-Gastaut syndrome (LGS): 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCOMETRIQDrug NamesCOMETRIQ

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Non-small cell lung cancer (NSCLC), differentiated thyroid carcinoma: papillary,

follicular, and Hurthle cell.

Exclusion Criteria

**Required Medical Information** For NSCLC: The requested medication is used for NSCLC when the patient's disease

expresses rearranged during transfection (RET) gene rearrangements.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group COPIKTRA
Drug Names COPIKTRA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Hepatosplenic T-Cell lymphoma, breast implant-associated anaplastic large cell

lymphoma (ALCL), peripheral T-Cell lymphoma

Exclusion Criteria -

Required Medical Information For chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL), breast

implant-associated anaplastic large cell lymphoma (ALCL), and peripheral T-Cell lymphoma: the patient has relapsed or refractory disease. For hepatosplenic T-Cell

lymphoma: the patient has refractory disease.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group CORTROPHIN Drug Names CORTROPHIN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information

For the following diagnoses, patient has experienced an inadequate treatment response to a parenteral or an oral glucocorticoid (for ophthalmic diseases only, inadequate response to a trial of a topical ophthalmic glucocorticoid is also acceptable):

1) For rheumatic disorders (e.g., psoriatic arthritis, rheumatoid arthritis, ankylosing spondylitis, acute gouty arthritis): The requested drug must be used as adjunctive

spondylitis, acute gouty arthritis): The requested drug must be used as adjunctive treatment, 2) For nephrotic syndrome: the requested drug must be requested for induction of diuresis or for remission of proteinuria, 3) For multiple sclerosis (MS): patient has an acute exacerbation of MS, 4) Collagen diseases (e.g., systemic lupus erythematosus, dermatomyositis, or polymyositis), 5) Dermatologic diseases (e.g., severe erythema multiforme, Stevens-Johnson syndrome, severe psoriasis), 6) Ophthalmic diseases, acute or chronic (e.g., iritis, keratitis, optic neuritis), 7)

Symptomatic sarcoidosis, 8) Allergic states (e.g., serum sickness, atopic dermatitis).

Age Restrictions -

Prescriber Restrictions -

**Coverage Duration** MS exacerbation: 3 wks. Allergic states: 1 month. All other diagnoses: 3 months

-

Prior Authorization Group COTELLIC Drug Names COTELLIC

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Central nervous system (CNS) cancer (i.e., glioma, glioblastoma, astrocytoma,

oligodendroglioma), adjuvant systemic therapy for cutaneous melanoma.

Exclusion Criteria -

**Required Medical Information** For central nervous system (CNS) cancer (i.e., glioma, glioblastoma, astrocytoma,

oligodendroglioma): 1) The tumor is positive for BRAF V600E activating mutation, AND 2) The requested drug will be used in combination with vemurafenib. For melanoma: 1) The tumor is positive for BRAF V600 activating mutation (e.g., V600E or V600K), AND 2) The requested drug will be used in combination with vemurafenib. AND 3) The

requested drug will be used for either of the following: a) unresectable, limited

resectable, or metastatic disease, b) adjuvant systemic therapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group CRINONE Drug Names CRINONE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Prophylaxis for premature birth in women with a short cervix

**Exclusion Criteria** Prescribed to promote fertility

Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCUVRIORDrug NamesCUVRIOR

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** Documented diagnosis of Wilson's disease in patients with stable disease who are

de-coppered and tolerant to penicillamine.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCYSTADROPSDrug NamesCYSTADROPS

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For cystinosis: 1) Diagnosis was confirmed by ANY of the following: a) the presence of

increased cystine concentration in leukocytes, OR b) genetic testing, OR c)

demonstration of corneal cystine crystals by slit lamp examination, AND 2) the patient

has corneal cystine crystal accumulation.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupCYSTAGONDrug NamesCYSTAGON

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For nephropathic cystinosis: Diagnosis was confirmed by ANY of the following: 1) the

presence of increased cystine concentration in leukocytes, OR 2) genetic testing, OR 3)

demonstration of corneal cystine crystals by slit lamp examination.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group CYSTARAN
Drug Names CYSTARAN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For cystinosis: 1) Diagnosis was confirmed by ANY of the following: a) the presence of

increased cystine concentration in leukocytes, OR b) genetic testing, OR c)

demonstration of corneal cystine crystals by slit lamp examination, AND 2) the patient

has corneal cystine crystal accumulation.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupDALFAMPRIDINEDrug NamesDALFAMPRIDINE ER

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For multiple sclerosis, patient must meet the following: For new starts, prior to initiating

therapy, patient demonstrates sustained walking impairment. For continuation of therapy: patient must have experienced an improvement in walking speed OR other

objective measure of walking ability since starting the requested drug.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group DAURISMO
Drug Names DAURISMO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Post induction therapy following response to previous therapy with the same regimen

for acute myeloid leukemia (AML). Relapsed/refractory AML as a component of

repeating the initial successful induction regimen.

Exclusion Criteria -

**Required Medical Information** For acute myeloid leukemia: 1) the requested drug must be used in combination with

cytarabine, 2) the patient is 75 years of age or older OR has comorbidities that preclude

intensive chemotherapy, AND 3) the requested drug will be used as treatment for induction therapy, post-induction therapy, or relapsed or refractory disease.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupDAYBUEDrug NamesDAYBUE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information -

**Age Restrictions** 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupDEFERASIROXDrug NamesDEFERASIROX

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For chronic iron overload due to blood transfusions: pretreatment serum ferritin level is

greater than 1000 mcg/L.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupDIACOMITDrug NamesDIACOMIT

PA Indication Indicator All FDA-approved Indications

Off-label Uses -Exclusion Criteria -Required Medical Information -

**Age Restrictions** 6 months of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group DICLOFENAC EPOLAMINE PATCH

Drug NamesDICLOFENAC EPOLAMINEPA Indication IndicatorAll FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information Must have a documented diagnosis of acute pain due to one of the following: minor

strain, sprain, or contusion.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupDOPTELETDrug NamesDOPTELET

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For thrombocytopenia in patients with chronic liver disease: Untransfused platelet count

prior to a scheduled procedure is less than 50,000/mcL. For chronic immune thrombocytopenia (ITP): 1) For new starts: a) Patient has had an inadequate response or is intolerant to prior therapy such as corticosteroids or immunoglobulins, AND b) Untransfused platelet count at any point prior to the initiation of the requested medication is less than 30,000/mcL OR 30,000 to 50,000/mcL with symptomatic bleeding or risk factor(s) for bleeding (e.g., undergoing a medical or dental procedure where blood loss is anticipated, comorbidities such as peptic ulcer disease and hypertension, anticoagulation therapy, profession or lifestyle that predisposes patient to trauma). 2) For continuation of therapy, platelet count response to the requested drug: a) Current platelet count is less than or equal to 200,000/mcL OR b) Current platelet count is greater than 200,000/mcL and less than or equal to 400,000/mcL and dosing

will be adjusted to a platelet count sufficient to avoid clinically important bleeding.

Age Restrictions 18 years of age or older

Coverage Duration Chronic liver disease: 1 month, ITP initial: 6 months, ITP reauthorization: Plan Year

Other Criteria

**Prescriber Restrictions** 

**Prior Authorization Group Drug Names** 

PA Indication Indicator

Off-label Uses **Exclusion Criteria Required Medical Information**  DUPIXENT DUPIXENT

All FDA-approved Indications

For atopic dermatitis (AD), initial therapy: 1) Patient has moderate-to-severe disease, AND 2) Patient has had an inadequate treatment response to either a topical corticosteroid or a topical calcineurin inhibitor. OR topical corticosteroids and topical calcineurin inhibitors are not advisable for the patient. For AD, continuation of therapy: Patient achieved or maintained positive clinical response. For oral corticosteroid dependent asthma, initial therapy: Patient has inadequate asthma control despite current treatment with both of the following medications: 1) High-dose inhaled corticosteroid AND 2) Additional controller (i.e., long acting beta2-agonist, long-acting, muscarinic antagonist, leukotriene modifier, or sustained-release theophylline) unless patient has an intolerance or contraindication to such therapies. For moderate-to-severe asthma, initial therapy: Patient has a baseline blood eosinophil count of at least 150 cells per microliter and their asthma remains inadequately controlled despite current treatment with both of the following medications: 1) Medium-to-high-dose inhaled corticosteroid, AND 2) Additional controller (i.e., long acting beta2-agonist, long-acting muscarinic antagonist, leukotriene modifier, or sustained-release theophylline) unless patient has an intolerance or contraindication to such therapies. For asthma, continuation of therapy: Asthma control has improved on treatment with the requested drug, as demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations or a reduction in the daily maintenance oral corticosteroid dose. For chronic rhinosinusitis with nasal polyposis (CRSwNP): 1) The requested drug is used as add-on maintenance treatment, AND 2) Patient has experienced an inadequate treatment response to Xhance (fluticasone). Atopic Dermatitis: 6 months of age or older, Asthma: 6 years of age or older, Chronic Rhinosinusitis with Nasal Polyposis and Prurigo Nodularis: 18 years of age or older,

Age Restrictions

**Prescriber Restrictions Coverage Duration** Other Criteria

AD, initial: 4 months, PN, initial: 6 months, All others: Plan Year

Eosinophilic Esophagitis: 1 year of age or older

For eosinophilic esophagitis (EoE), initial therapy: 1) Diagnosis has been confirmed by esophageal biopsy, AND 2) Patient weighs at least 15 kilograms, AND 3) Patient experienced an inadequate treatment response, intolerance, or patient has a contraindication to a topical corticosteroid (e.g., fluticasone propionate or budesonide). For EoE, continuation of therapy: Patient achieved or maintained a positive clinical response. For prurigo nodularis (PN), initial therapy: Patient has had an inadequate treatment response to a topical corticosteroid OR topical corticosteroids are not advisable for the patient. For PN, continuation of therapy: Patient achieved or maintained a positive clinical response.

Updated 07/01/2024 28 **Prior Authorization Group** EGRIFTA **Drug Names** EGRIFTA SV

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

**Exclusion Criteria** Use for weight loss

**Required Medical Information** For human immunodeficiency virus (HIV)-infected patients with lipodystrophy: Patient is

receiving anti-retroviral therapy. For patients who have received at least 6 months of the requested drug: Patient has demonstrated clear clinical improvement from baseline that is supported by a waist circumference measurement or computed tomography (CT)

scan.

Age Restrictions

Prescriber Restrictions Prescribed by or in consultation with an infectious disease specialist or endocrinologist

Coverage Duration 6 months
Other Criteria -

Prior Authorization Group ELIGARD
Drug Names ELIGARD

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent androgen receptor positive salivary gland tumors

Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** EMFLAZA

Drug NamesDEFLAZACORT, EMFLAZAPA Indication IndicatorAll FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of Duchenne muscular dystrophy (DMD).

Age Restrictions -

**Prescriber Restrictions**Must be prescribed by a neurologist or a provider who specializes in the treatment of

DMD.

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** 

**Drug Names** 

**PA Indication Indicator** 

Off-label Uses

Exclusion Criteria
Required Medical Information

**ENBREL** 

ENBREL, ENBREL MINI, ENBREL SURECLICK

All FDA-approved Indications, Some Medically-accepted Indications Hidradenitis suppurativa, non-radiographic axial spondyloarthritis

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For moderately to severely active rheumatoid arthritis (new starts only): 1) patient has experienced an inadequate treatment response, intolerance, or has a contraindication to methotrexate (MTX) OR 2) patient has experienced an inadequate treatment response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD. For active ankylosing spondylitis and non-radiographic axial spondyloarthritis (new starts only): patient has experienced an inadequate treatment response or intolerance to a non-steroidal anti-inflammatory drug (NSAID) OR the patient has a contraindication that would prohibit a trial of NSAIDs. For moderate to severe plaque psoriasis (new starts only): at least 3% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin, intertriginous areas) are affected at the time of diagnosis. For hidradenitis suppurativa (new starts only): patient has severe, refractory disease.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupENSPRYNGDrug NamesENSPRYNG

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupEPCLUSADrug NamesEPCLUSA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For hepatitis C virus (HCV): Infection confirmed by presence of HCV RNA in the serum

prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of human immunodeficiency virus (HIV) coinfection, presence or absence of resistance-associated substitutions where applicable, transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current American Association for the Study of Liver Diseases and Infectious Diseases Society of America (AASLD-IDSA) treatment

guidelines.

Age Restrictions Prescriber Restrictions -

Coverage Duration Criteria will be applied consistent with current AASLD-IDSA guidance

Other Criteria -

Prior Authorization GroupEPIDIOLEXDrug NamesEPIDIOLEX

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information -

**Age Restrictions** 1 year of age or older

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupEPRONTIADrug NamesEPRONTIA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For treatment of partial-onset seizures (i.e., focal-onset seizures): 1)The patient has

experienced an inadequate treatment response, intolerance, or has a contraindication to a generic anticonvulsant AND 2) the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to any of the following:

Aptiom (if 4 years of age or older), Xcopri (if 18 years of age or older), Spritam (if 4 years of age or older). For monotherapy treatment of primary generalized tonic-clonic seizures: 1) The patient has experienced an inadequate treatment response or intolerance to topiramate tablets or capsules, OR 2) The patient has difficulty swallowing solid oral dosage forms (e.g., tablets, capsules). For adjunctive treatment of primary generalized tonic-clonic seizures: 1) The patient has experienced an inadequate treatment response, intolerance, or has a contraindication to a generic anticonvulsant AND 2) If the patient is 6 years of age or older, the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to Spritam. For the preventative treatment of migraines: 1) The patient has experienced an inadequate treatment response or intolerance to topiramate tablets or capsules, OR 2) The patient has difficulty swallowing solid oral dosage forms (e.g., tablets, capsules).

**Age Restrictions** Epilepsy: 2 years of age or older, Migraine: 12 years of age or older **Prescriber Restrictions** -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** ERGOTAMINE

Drug Names ERGOTAMINE TARTRATE/CAFFE

PA Indication Indicator All FDA-approved Indications
Off-label Uses -

Exclusion Criteria Coverage will be denied when used in conjunction with potent CYP3A4 inhibitors (e.g.,

ritonavir, nelfinavir, indinavir, erythromycin, clarithromycin).

**Required Medical Information** The patient has experienced an inadequate treatment response, intolerance, or has a

contraindication to at least ONE triptan 5-HT1 agonist.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupERIVEDGEDrug NamesERIVEDGE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Adult medulloblastoma

Exclusion Criteria -

**Required Medical Information** For adult medulloblastoma: patient has received prior systemic therapy AND has

tumor(s) with mutations in the sonic hedgehog pathway.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ERLEADA
Drug Names ERLEADA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** The requested drug will be used in combination with a gonadotropin-releasing hormone

(GnRH) analog or after bilateral orchiectomy.

Age Restrictions --

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** ERLOTINIB

**Drug Names** ERLOTINIB HYDROCHLORIDE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent non-small cell lung cancer (NSCLC), recurrent chordoma, relapsed or stage

IV renal cell carcinoma (RCC), brain metastases from non-small cell lung cancer

(NSCLC), recurrent pancreatic cancer.

Exclusion Criteria -

**Required Medical Information** For NSCLC (including brain metastases from NSCLC): 1) the disease is recurrent,

advanced, or metastatic and 2) the patient has sensitizing EGFR mutation-positive disease. For pancreatic cancer: the disease is locally advanced, unresectable,

recurrent, or metastatic.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ESBRIET
Drug Names PIRFENIDONE

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For idiopathic pulmonary fibrosis (new starts only): 1) a high-resolution computed

tomography (HRCT) study of the chest or a lung biopsy reveals the usual interstitial pneumonia (UIP) pattern, OR 2) HRCT study of the chest reveals a result other than the UIP pattern (e.g., probable UIP, indeterminate for UIP) and the diagnosis is supported either by a lung biopsy or by a multidisciplinary discussion between at least a radiologist and pulmonologist who are experienced in idiopathic pulmonary fibrosis if a

lung biopsy has not been conducted.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupEUCRISADrug NamesEUCRISA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information

For mild to moderate atopic dermatitis, the patient meets either of the following criteria:

1) If the patient is 2 years of age or older and the requested drug will be used on sensitive skin areas (e.g., face, genitals, or skin folds), the patient experienced an inadequate treatment response, intolerance, or contraindication to a topical calcineurin inhibitor OR 2) If the patient is 2 years of age or older and the requested drug is being prescribed for use on non-sensitive (or remaining) skin areas, the patient experienced an inadequate treatment response, intolerance, or contraindication to a medium or higher potency topical corticosteroid or a topical calcineurin inhibitor.

Age Restrictions 3 months of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group EVENITY
Drug Names EVENITY

PA Indication Indicator All FDA-ar

Off-label Uses
Exclusion Criteria

Required Medical Information

All FDA-approved Indications

Patients who have had a myocardial infarction or stroke within the preceding year. For postmenopausal osteoporosis, patient has ONE of the following: 1) history of fragility fracture, OR 2) pre-treatment T-score of less than or equal to -2.5 or pre-treatment T-score greater than -2.5 and less than -1 with a high pre-treatment Fracture Risk Assessment Tool (FRAX) fracture probability AND patient has ANY of the following: a) indicators for higher fracture risk (e.g., advanced age, frailty, glucocorticoid therapy, very low T-scores, or increased fall risk), or b) patient has failed prior treatment with or is intolerant to a previous injectable osteoporosis therapy, or c) patient has had an oral bisphosphonate trial of at least 1-year duration or there is a clinical reason to avoid treatment with an oral bisphosphonate.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

12 months lifetime total

Patient has high Fracture Risk Assessment Tool (FRAX) fracture probability if the 10 year probability is either greater than or equal to 20 percent for any major osteoporotic fracture or greater than or equal to 3 percent for hip fracture. The estimated risk score generated with FRAX should be multiplied by 1.15 for major osteoporotic fracture and 1.2 for hip fracture if glucocorticoid treatment is greater than 7.5 mg (prednisone equivalent) per day.

**Prior Authorization Group Drug Names** PA Indication Indicator Off-label Uses

**EVEROLIMUS EVEROLIMUS** 

All FDA-approved Indications, Some Medically-accepted Indications Classic Hodgkin lymphoma, thymomas and thymic carcinomas, previously treated Waldenstrom's macroglobulinemia/lymphoplasmacytic lymphoma, soft tissue sarcoma (perivascular epithelioid cell tumors (PEComa) and lymphangioleiomyomatosis subtypes), gastrointestinal stromal tumors, neuroendocrine tumors of the thymus, well differentiated grade 3 neuroendocrine tumors, thyroid carcinoma (papillary, Hurthle cell, and follicular), endometrial carcinoma, histiocytic neoplasms (Rosai-Dorfman Disease, Erdheim-Chester Disease, Langerhans Cell Histiocytosis)

**Exclusion Criteria Required Medical Information** 

For breast cancer: 1) The disease is recurrent unresectable, advanced, or metastatic hormone receptor (HR)-positive, human epidermal growth factor receptor 2 (HER2)-negative, AND 2) The requested drug is prescribed in combination with exemestane, fulvestrant, or tamoxifen, AND 3) The requested drug is used for subsequent treatment. For renal cell carcinoma: The disease is relapsed, advanced, or stage IV. For subependymal giant cell astrocytoma (SEGA): The requested drug is given as adjuvant treatment. For gastrointestinal stromal tumor: The disease is recurrent/progressive, unresectable, or metastatic AND the patient failed an FDA-approved therapy (e.g., imatinib, sunitinib, regorafenib, ripretinib). For symptomatic or relapsed/refractory Erdheim-Chester Disease (ECD), symptomatic or relapsed/refractory Rosai-Dorfman Disease, and Langerhans Cell Histiocytosis (LCH): the patient must have a phosphatidylinositol-4,5-bisphosphate 3-kinase catalytic subunit alpha (PIK3CA) mutation.

Age Restrictions **Prescriber Restrictions Coverage Duration** Other Criteria

Plan Year

Updated 07/01/2024 36 Prior Authorization GroupEVRYSDIDrug NamesEVRYSDI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For spinal muscular atrophy (SMA) initial therapy, patient meets all of the following: 1)

Patient has type 1, type 2, or type 3 SMA, and 2) Patient is not dependent on permanent ventilation. For SMA continuation of therapy, patient meets all of the

following: 1) Patient has type 1, type 2, or type 3 SMA, AND 2) Patient has experienced clinically significant functional improvement or maintenance of muscle function.

Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with a physician who specializes in spinal muscular

atrophy

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** EXKIVITY **Drug Names** EXKIVITY

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupFABHALTADrug NamesFABHALTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For paroxysmal nocturnal hemoglobinuria (PNH) (initial): 1) the diagnosis of PNH was

confirmed by detecting a deficiency of glycosylphosphatidylinositol-anchored proteins (GPI-APs) AND 2) flow cytometry is used to demonstrate GPI-AP deficiency. For PNH (continuation): 1) there is no evidence of unacceptable toxicity or disease progression while on the current regimen AND 2) the patient has demonstrated a positive response

to therapy.

Age Restrictions -

Prescriber Restrictions -

**Coverage Duration** Initial: 6 months, Continuation: Plan Year

Other Criteria -

**Prior Authorization Group** FABRAZYME **Drug Names** FABRAZYME

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For Fabry disease, the patient meets ANY of the following: 1) diagnosis of Fabry

disease was confirmed by an enzyme assay demonstrating a deficiency of alpha-galactosidase enzyme activity or by genetic testing,  $\mathsf{OR}\ \mathsf{2}$ ) the patient is a

symptomatic obligate carrier.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** FASENRA

Drug NamesFASENRA, FASENRA PENPA Indication IndicatorAll FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Severe asthma, initial therapy: 1) Either a) Patient has baseline blood eosinophil count

of at least 150 cells per microliter OR b) Patient is dependent on systemic

corticosteroids, and 2) Patient has a history of severe asthma despite current treatment with both of the following medications: a) medium-to-high-dose inhaled corticosteroid and b) additional controller (i.e., long-acting beta2-agonist, long-acting muscarinic antagonist, leukotriene modifier, or sustained-release theophylline) unless patient has an intolerance or contraindication to such therapies. Severe asthma, continuation of therapy: Asthma control has improved on treatment with the requested drug, as demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations or a reduction in the daily maintenance oral corticosteroid dose.

Age Restrictions 6 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** FENTANYL PATCH

**Drug Names** FENTANYL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** The requested drug is being prescribed for pain associated with cancer, sickle cell

disease, a terminal condition, or pain being managed through palliative care OR the patient meets all of the following: 1) The requested drug is being prescribed for pain severe enough to require daily, around-the-clock, long-term treatment in a patient who has been taking an opioid AND 2) The patient can safely take the requested dose based on their history of opioid use [Note: This drug should be prescribed only by healthcare professionals who are knowledgeable in the use of potent opioids for the management of chronic pain.] AND 3) The patient has been evaluated and the patient will be monitored for the development of opioid use disorder AND 4) This request is for continuation of therapy for a patient who has been receiving an extended-release opioid agent for at least 30 days OR the patient has taken an immediate-release opioid for at least one week.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group FILSPARI
Drug Names FILSPARI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information

For reduction of proteinuria in patients with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression: 1) The patient had an inadequate response to therapy with a maximally tolerated dose of a renin-angiotensin system (RAS) inhibitor (e.g., angiotensin-converting enzyme [ACE] inhibitor or angiotensin-receptor blocker [ARB]) OR 2) The patient experienced an intolerance or has a contraindication to RAS inhibitors.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupFINTEPLADrug NamesFINTEPLA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information -

**Age Restrictions** 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupFIRMAGONDrug NamesFIRMAGON

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of advanced Prostate cancer.

Age Restrictions -

**Prescriber Restrictions**Must be prescribed by an oncologist or a urologist specializing in prostate cancer.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupFOTIVDADrug NamesFOTIVDA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For advanced renal cell carcinoma: the following criteria must be met: 1) The disease is

relapsed or refractory, 2) The requested drug must be used after at least two prior systemic therapies, and 3) The patient has experienced disease progression or an

intolerable adverse event with a trial of Cabometyx (cabozantinib).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupFRUZAQLADrug NamesFRUZAQLA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupFYCOMPADrug NamesFYCOMPA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For treatment of partial-onset seizures (i.e., focal-onset seizures): 1) The patient has

experienced an inadequate treatment response, intolerance, or has a contraindication to a generic anticonvulsant AND 2) The patient has experienced an inadequate treatment response, intolerance, or has a contraindication to any of the following:

Aptiom, Xcopri, Spritam. For adjunctive treatment of primary generalized tonic-clonic

seizures: 1) The patient has experienced an inadequate treatment response,

intolerance, or has a contraindication to a generic anticonvulsant AND 2) The patient

has experienced an inadequate treatment response, intolerance, or has a

contraindication to Spritam.

**Age Restrictions** Partial-onset seizures (i.e., focal-onset seizures): 4 years of age or older. Primary

generalized tonic-clonic seizures: 12 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** GATTEX **Drug Names** GATTEX

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For short bowel syndrome (SBS) initial therapy: 1) If the request is for an adult patient,

the patient has been dependent on parenteral support for at least 12 months OR 2) If the request is for a pediatric patient, the patient is dependent on parenteral support. For SBS continuation: Requirement for parenteral support has decreased from baseline

while on therapy with the requested drug.

Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with a gastroenterologist, gastrointestinal surgeon, or

nutritional support specialist.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupGAVRETODrug NamesGAVRETO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent rearranged during transfection (RET) rearrangement-positive non-small cell

lung cancer

Exclusion Criteria -

**Required Medical Information** For non-small cell lung cancer, patient must meet all of the following: 1) The disease is

recurrent, advanced, or metastatic, and 2) The tumor is rearranged during transfection

(RET) fusion-positive or RET rearrangement-positive.

Age Restrictions Non-small cell lung cancer: 18 years of age or older. Medullary thyroid cancer and

thyroid cancer: 12 years of age or older.

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** GILENYA

**Drug Names** FINGOLIMOD HYDROCHLORIDE, GILENYA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -Required Medical Information -

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupGILOTRIFDrug NamesGILOTRIF

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For non-small cell lung cancer (NSCLC): Patient meets either of the following: 1)

Patient has metastatic squamous NSCLC that progressed after platinum-based

chemotherapy, OR 2) Patient has sensitizing epidermal growth factor receptor (EGFR)

mutation-positive disease.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** GLATIRAMER

**Drug Names** GLATIRAMER ACETATE, GLATOPA

PA Indication Indicator All FDA-approved Indications

Off-label Uses
Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupGOCOVRIDrug NamesGOCOVRI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** GRALISE

**Drug Names** GABAPENTIN ONCE-DAILY, GRALISE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Postherpetic neuralgia: The patient has experienced an inadequate treatment response

or intolerance to gabapentin immediate-release

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group Drug Names

PA Indication Indicator
Off-label Uses
Exclusion Criteria
Required Medical Information

Age Restrictions
Prescriber Restrictions

Coverage Duration
Other Criteria

**GROWTH HORMONE** 

GENOTROPIN, GENOTROPIN MINIQUICK, HUMATROPE, NORDITROPIN FLEXPRO, NUTROPIN AQ NUSPIN 10, NUTROPIN AQ NUSPIN 20, NUTROPIN AQ NUSPIN 5, OMNITROPE, SOGROYA, ZOMACTON

All Medically-accepted Indications

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Pediatric patients with closed epiphyses

Pediatric growth hormone deficiency (GHD): Patient (pt) is a neonate or was diagnosed with GHD as a neonate OR meets any of the following: 1) younger than 2.5 years old (yo) with pre-treatment (pre-tx) height (ht) more than 2 standard deviations (SD) below mean and slow growth velocity OR 2) 2.5 yo or older AND one of the following: a) pre-tx 1-year ht velocity more than 2 SD below mean OR b) pre-tx ht more than 2 SD below mean and 1-year ht velocity more than 1 SD below mean, AND patient meets any of the following: 1) failed 2 pre-tx growth hormone (GH) stimulation tests (peak below 10 ng/mL), OR 2) pituitary/central nervous system (CNS) disorder (e.g., genetic defects, acquired structural abnormalities, congenital structural abnormalities) and pre-tx insulin-like growth factor-1 (IGF-1) more than 2 SD below mean. Turner syndrome (TS): 1) Confirmed by karyotyping AND 2) pre-tx ht is less than the 5th percentile for age. Small for gestational age (SGA): 1) Birth weight (wt) less than 2500g at gestational age (GA) greater than 37 weeks, OR birth wt or length below 3rd percentile for GA or at least 2 SD below mean for GA, AND 2) did not manifest catch-up growth by age 2.

SGA: 2 years of age or older

Prescribed by or in consultation with an endocrinologist, nephrologist, infectious disease specialist, gastroenterologist/nutritional support specialist, or geneticist. Plan Year

Adult GHD: Pt meets any of the following: 1) failed 2 pre-tx GH stimulation tests, OR 2) pre-tx IGF-1 more than 2 SD below mean AND failed 1 pre-tx GH stimulation test. (Note: Stimulation tests include: a) insulin tolerance test [ITT] [peak GH less than or equal to 5 ng/ml], or b) Macrilen-stimulation test [peak GH level less than 2.8ng/ml], or c) glucagon-stimulation test [GST] [peak GH level less than or equal to 3 ng/ml] for pt with a body mass index [BMI] 25-30 kg/m2 and high pretest probability of GHD [e.g., acquired structural abnormalities] or BMI less than 25 kg/m2, or d) GST [peak GH level less than or equal to 1 ng/ml] in pt with BMI 25-30 kg/m2 and low pretest probability of GHD or BMI greater than 30 kg/m2), OR 3) organic hypothalamic-pituitary disease (e.g., suprasellar mass with previous surgery and cranial irradiation) with 3 or more pituitary hormone deficiencies AND pre-tx IGF-1 more than 2 SD below mean, OR 4) genetic or structural hypothalamic-pituitary defects, OR 5) childhood-onset GHD with congenital (genetic or structural) abnormality of the hypothalamus/pituitary/CNS. Renewal for pediatric GHD, TS, SGA, and adult GHD: Patient is experiencing improvement.

**Prior Authorization Group HAEGARDA Drug Names HAEGARDA** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** 

For the prevention of acute angioedema attacks due to hereditary angioedema (HAE): The patient meets either of the following: 1) the patient has hereditary angioedema (HAE) with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing OR 2) the patient has hereditary angioedema with normal C1 inhibitor confirmed by laboratory testing and either of the following: a) patient tested positive for an F12, angiopoietin-1. plasminogen, kininogen-1 (KNG1), heparan sulfate-glucosamine 3-O-sulfotransferase 6 (HS3ST6), or myoferlin (MYOF) gene mutation OR b) patient has a family history of angioedema and the angioedema was refractory to a trial of high-dose antihistamine therapy for at least one month.

Age Restrictions 6 years of age or older

**Prescriber Restrictions** Prescribed by or in consultation with an immunologist, allergist, or rheumatologist

**Coverage Duration** Plan Year

Other Criteria

**Prior Authorization Group** HARVONI **Drug Names HARVONI** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** 

For hepatitis C virus (HCV): Infection confirmed by presence of HCV RNA in the serum prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of human immunodeficiency virus (HIV) coinfection, presence or absence of resistance-associated substitutions where applicable, transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current American Association for the Study of Liver Diseases and Infectious Diseases Society of America (AASLD-IDSA) treatment quidelines.

Age Restrictions **Prescriber Restrictions** 

**Coverage Duration** 

Criteria applied consistent w/ current AASLD-IDSA guidance. Reminder for 8wk option

if appropriate.

Other Criteria

Updated 07/01/2024 46 Prior Authorization Group Drug Names PA Indication Indicator Off-label Uses HERCEPTIN HERCEPTIN HYLECTA

All FDA-approved Indications, Some Medically-accepted Indications

Neoadjuvant treatment for human epidermal growth factor receptor 2 (HER2)-positive breast cancer, recurrent or advanced unresectable HER2-positive breast cancer, leptomeningeal metastases from HER2-positive breast cancer, brain metastases from HER2-positive breast cancer, hereast cancer, recurrent, or metastatic uterine serous carcinoma, hereast cancer, and breast and breast allowed cancer (including appendiceal adenocarcinoma), hereast and breast allowed carcinoma (gallbladder cancer, intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma), hereast cancer, intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma, extr

Exclusion Criteria
Required Medical Information

All indications: the patient had an intolerable adverse event to Trazimera and that adverse event was NOT attributed to the active ingredient as described in the prescribing information. For colorectal cancer (including appendiceal adenocarcinoma):

1) the disease is HED2 amplified and DAS and DDAE wild type and 2) the requested.

1) the disease is HER2-amplified and RAS and BRAF wild-type and 2) the requested drug is used in combination with pertuzumab, tucatinib or lapatinib and 3) the patient has not had previous treatment with a HER2 inhibitor. For hepatobiliary carcinoma: 1) the disease is HER2-positive AND 2) the requested drug is used in combination with

pertuzumab.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization Group Drug Names PA Indication Indicator Off-label Uses HERZUMA HERZUMA

All FDA-approved Indications, Some Medically-accepted Indications
Neoadjuvant treatment for human epidermal growth factor receptor 2 (HER2)-positive
breast cancer, recurrent or advanced unresectable HER2-positive breast cancer,
leptomeningeal metastases from HER2-positive breast cancer, brain metastases from
HER2-positive breast cancer, HER2-positive esophageal and esophagogastric junction
adenocarcinoma, HER2-positive advanced, recurrent, or metastatic uterine serous
carcinoma, HER2-amplified and RAS and BRAF wild-type colorectal cancer (including
appendiceal adenocarcinoma), HER2-positive recurrent salivary gland tumor,
HER2-positive unresectable or metastatic hepatobiliary carcinoma (gallbladder cancer,
intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma), HER2
overexpression positive locally advanced, unresectable, or recurrent gastric
adenocarcinoma.

Exclusion Criteria
Required Medical Information

All indications: the patient had an intolerable adverse event to Trazimera and that adverse event was NOT attributed to the active ingredient as described in the prescribing information. For colorectal cancer (including appendiceal adenocarcinoma): 1) the disease is HER2-amplified and RAS and BRAF wild-type and 2) the requested drug is used in combination with pertuzumab, tucatinib or lapatinib and 3) the patient has not had previous treatment with a HER2 inhibitor. For hepatobiliary carcinoma: 1) the disease is HER2-positive AND 2) the requested drug is used in combination with pertuzumab.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** HETLIOZ

**Drug Names** TASIMELTEON

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For Non-24-Hour Sleep-Wake Disorder: 1) For initial therapy and continuation of

therapy the patient must meet both of the following: a) diagnosis of total blindness in both eyes (e.g., nonfunctioning retinas) and b) unable to perceive light in either eye, AND 2) If currently on therapy with the requested drug, patient must meet at least one of the following: a) increased total nighttime sleep or b) decreased daytime nap duration. For nighttime sleep disturbances in Smith-Magenis Syndrome (SMS): 1) For initial therapy and continuation therapy, the patient has a confirmed diagnosis of SMS AND 2) If currently on therapy with the requested drug, the patient experienced

improvement in the quality of sleep since starting therapy.

Age Restrictions Non-24: 18 years of age or older. SMS: 16 years of age or older

**Prescriber Restrictions** Prescribed by or in consultation with a sleep disorder specialist, neurologist, or

psychiatrist.

**Coverage Duration** Initiation: 6 Months, Renewal: Plan Year

Other Criteria -

Prior Authorization GroupHETLIOZ LQDrug NamesHETLIOZ LQ

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For nighttime sleep disturbances in Smith-Magenis Syndrome (SMS): 1) For initial

therapy and continuation therapy, the patient has a confirmed diagnosis of SMS AND 2) If currently on therapy with the requested drug, the patient experienced improvement

in the quality of sleep since starting therapy.

Age Restrictions 3 to 15 years of age

Prescriber Restrictions Prescribed by or in consultation with a sleep disorder specialist or neurologist

**Coverage Duration** Initiation: 6 Months, Renewal: Plan Year.

Other Criteria -

Prior Authorization Group Drug Names

**HUMIRA** 

ADALIMUMAB-AACF (2 PEN), HADLIMA, HADLIMA PUSHTOUCH, HUMIRA,

HUMIRA PEDIATRIC CROHNS D, HUMIRA PEN, HUMIRA PEN-CD/UC/HS START, HUMIRA PEN-PEDIATRIC UC S, HUMIRA PEN-PS/UV STARTER, IDACIO (2 PEN),

IDACIO (2 SYRINGE), IDACIO STARTER PACKAGE FO

PA Indication Indicator

All Medically-accepted Indications

Off-label Uses
Exclusion Criteria

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**Required Medical Information** 

For moderately to severely active rheumatoid arthritis (new starts only): 1) patient has experienced an inadequate treatment response, intolerance, or has a contraindication to methotrexate (MTX) OR 2) patient has experienced an inadequate treatment response or intolerance to a prior biologic disease-modifying antirheumatic drug

response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD. For active ankylosing spondylitis and non-radiographic axial spondyloarthritis (new starts only): patient has experienced an inadequate treatment response or intolerance to a non-steroidal anti-inflammatory drug (NSAID) OR the patient has a contraindication that would prohibit a trial of NSAIDs. For moderate to severe plaque psoriasis (new starts only): at least 3% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin,

intertriginous areas) are affected at the time of diagnosis.

Age Restrictions

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**Prescriber Restrictions** 

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Coverage Duration
Other Criteria

Plan Year

For non-infectious intermediate, posterior and panuveitis (new starts only): 1) patient has experienced an inadequate treatment response or intolerance to a corticosteroid OR 2) the patient has a contraindication that would prohibit a trial of corticosteroids.

**Prior Authorization Group** 

IBRANCE

**Drug Names** 

**IBRANCE** 

**PA Indication Indicator** 

All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Unresectable well-differentiated/dedifferentiated liposarcoma of the retroperitoneum, recurrent hormone receptor-positive human epidermal growth factor receptor 2

(HER2)-negative breast cancer

**Exclusion Criteria** 

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**Required Medical Information** 

-

Age Restrictions

-

**Prescriber Restrictions** 

Plan Year

Coverage Duration
Other Criteria

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**Prior Authorization Group** ICATIBANT

**Drug Names**ICATIBANT ACETATE, SAJAZIR **PA Indication Indicator**All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For acute angioedema attacks due to hereditary angioedema (HAE): Patient meets

either of the following: 1) the patient has HAE with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing OR 2) the patient has HAE with normal C1 inhibitor confirmed by laboratory testing and one of the following: a) the patient tested positive

for an F12, angiopoietin-1, plasminogen, kininogen-1 (KNG1), heparan

sulfate-glucosamine 3-O-sulfotransferase 6 (HS3ST6), or myoferlin (MYOF) gene mutation OR b) the patient has a family history of angioedema and the angioedema was refractory to a trial of high-dose antihistamine therapy for at least one month.

Age Restrictions 18 years of age or older

Prescriber Restrictions Prescribed by or in consultation with an immunologist, allergist, or rheumatologist

Coverage Duration Plan Year
Other Criteria -

Prior Authorization Group ICLUSIG
Drug Names ICLUSIG

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Myeloid and/or lymphoid neoplasms with eosinophilia and FGFR1 or ABL1

rearrangement in the chronic phase or blast phase

Exclusion Criteria -

**Required Medical Information** For chronic myeloid leukemia (CML), including patients who have received a

hematopoietic stem cell transplant: 1) patient has accelerated or blast phase CML and no other kinase inhibitor is indicated OR 2) patient has chronic phase CML and has experienced resistance or intolerance to at least 2 prior kinase inhibitors AND at least one of those was imatinib or dasatinib OR 3) patient is positive for the T315I mutation. For acute lymphoblastic leukemia (ALL), including patients who have received a hematopoietic stem cell transplant: diagnosis was confirmed by detection of the

Philadelphia chromosome or BCR-ABL gene.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group IDHIFA
Drug Names IDHIFA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Newly-diagnosed acute myeloid leukemia

Exclusion Criteria -

**Required Medical Information** For acute myeloid leukemia (AML) with an isocitrate dehydrogenase-2 (IDH2) mutation:

1) patient is 60 years of age or older with newly-diagnosed AML and meets one of the following: a) patient is not a candidate for intensive induction therapy, or b) patient declines intensive induction chemotherapy, OR 2) patient is 60 years of age or older and the requested drug will be used as post-induction therapy following response to induction therapy with the requested drug. OR 3) patient has relapsed or refractory

AML.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** IMATINIB

**Drug Names** IMATINIB MESYLATE

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Desmoid tumors, pigmented villonodular synovitis/tenosynovial giant cell tumor

(PVNS/TGCT), recurrent chordoma, melanoma, Kaposi sarcoma, chronic graft versus

host disease (cGVHD), T-cell acute lymphoblastic leukemia with ABL-class translocation, aggressive systemic mastocytosis for well-differentiated systemic mastocytosis (WDSM) or when eosinophilia is present with FIP1L1-PDGFRA fusion

gene, myeloid and/or lymphoid neoplasms with eosinophilia and ABL1,

FIP1L1-PDGFRA, or PDGFRB rearrangement in the chronic phase or blast phase

Exclusion Criteria

**Required Medical Information** For chronic myeloid leukemia (CML) or Philadelphia chromosome positive acute

lymphoblastic leukemia (Ph+ ALL), including patients who have received a hematopoietic stem cell transplant: diagnosis was confirmed by detection of the Philadelphia chromosome or BCR-ABL gene. For CML: patient did not fail (excluding failure due to intolerance) prior therapy with a tyrosine kinase inhibitor. For melanoma:

c-Kit mutation is positive.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group Drug Names PA Indication Indicator Off-label Uses IMBRUVICA IMBRUVICA

All FDA-approved Indications, Some Medically-accepted Indications
Hairy cell leukemia, lymphoplasmacytic lymphoma, primary central nervous system
(CNS) lymphoma, Human Immunodeficiency Virus (HIV) -related B-cell lymphoma,
diffuse large B-cell lymphoma, post-transplant lymphoproliferative disorders, high-grade
B-cell lymphoma, mantle cell lymphoma, marginal zone lymphoma (including
extranodal marginal zone lymphoma of the stomach, extranodal marginal zone
lymphoma of nongastric sites, nodal marginal zone lymphoma, splenic marginal zone
lymphoma)

Exclusion Criteria
Required Medical Information

For mantle cell lymphoma: 1) the requested drug will be used as second-line or subsequent therapy, OR 2) the requested drug will be used in combination with rituximab as pretreatment to induction therapy with RHyperCVAD (rituximab, cyclophosphamide, vincristine, doxorubicin, and dexamethasone) regimen, OR 3) the requested drug will be used as aggressive induction therapy. For marginal zone lymphoma (including extranodal marginal zone lymphoma of the stomach, extranodal marginal zone lymphoma of nongastric sites, nodal marginal zone lymphoma, and splenic marginal zone lymphoma): the requested drug will be used as second-line or subsequent therapy. For hairy cell leukemia: the requested drug will be used as a single agent for disease progression. For primary CNS lymphoma: 1) the disease is relapsed or refractory, OR 2) the requested drug is used for induction therapy as a single agent. For diffuse large B-cell lymphoma and high-grade B-cell lymphoma: the requested drug will be used as second-line or subsequent therapy. For HIV-related B-cell lymphoma: the requested drug will be used as a single agent and as second-line or subsequent therapy for relapsed disease. For post-transplant lymphoproliferative disorders: the requested drug will be used in patients who have received prior chemoimmunotherapy.

Age Restrictions
Prescriber Restrictions
Coverage Duration

-

Plan Year

Other Criteria

Prior Authorization Group INCRELEX Drug Names INCRELEX

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

**Exclusion Criteria** Pediatric patients with closed epiphyses

**Required Medical Information** For growth failure due to severe primary insulin-like growth factor-1 (IGF-1) deficiency

or growth hormone (GH) gene deletion in patients who have developed neutralizing antibodies to GH, patient meets all of the following prior to beginning therapy with the requested drug (new starts only): 1) height 3 or more standard deviations (SD) below the mean for children of the same age and gender AND 2) basal IGF-1 level 3 or more SD below the mean for children of the same age and gender AND 3) provocative growth hormone test showing a normal or elevated growth hormone level. For growth failure due to severe primary IGF-1 deficiency or GH gene deletion in patients who have developed neutralizing antibodies to GH, continuation of therapy: patient is

experiencing improvement.

**Age Restrictions** 2 years of age or older

**Prescriber Restrictions** Prescribed by or in consultation with an endocrinologist

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group INGREZZA
Drug Names INGREZZA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of tardive dyskinesia or chorea associated with

Huntington's disease

Age Restrictions -

**Prescriber Restrictions**Must be prescribed by a neurologist or with consultation by a neurologist.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group INLYTA
Drug Names INLYTA

PA Indication Indicator

All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Thyroid carcinoma (papillary, Hurthle cell, or follicular), alveolar soft part sarcoma

Exclusion Criteria -

**Required Medical Information** For renal cell carcinoma: The disease is advanced, relapsed, or stage IV.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupINQOVIDrug NamesINQOVI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group INREBIC Drug Names INREBIC

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia and janus kinase 2

(JAK2) rearrangement, accelerated phase myelofibrosis, blast phase

myelofibrosis/acute myeloid leukemia

Exclusion Criteria

**Required Medical Information** For myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia and JAK2

rearrangement: the disease is in chronic or blast phase.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group Drug Names** 

IR BEFORE ER

All FDA-approved Indications

HYDROCODONE BITARTRATE ER. HYDROMORPHONE HCL ER.

HYDROMORPHONE HYDROCHLORI, METHADONE HCL, MORPHINE SULFATE ER, OXYCODONE HYDROCHLORIDE E, OXYMORPHONE HYDROCHLORIDE.

TRAMADOL HCL ER, TRAMADOL HYDROCHLORIDE ER

PA Indication Indicator

Off-label Uses

**Exclusion Criteria** 

**Required Medical Information** 

The requested drug is being prescribed for pain associated with cancer, sickle cell disease, a terminal condition, or pain being managed through palliative care OR the patient meets all of the following: 1) The requested drug is being prescribed for pain severe enough to require daily, around-the-clock, long-term treatment in a patient who has been taking an opioid AND 2) The patient can safely take the requested dose based on their history of opioid use [Note: This drug should be prescribed only by healthcare professionals who are knowledgeable in the use of potent opioids for the management of chronic pain.] AND 3) The patient has been evaluated and the patient will be monitored for the development of opioid use disorder AND 4) This request is for continuation of therapy for a patient who has been receiving an extended-release opioid agent for at least 30 days OR the patient has taken an immediate-release opioid for at least one week.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** Plan Year

Other Criteria

**Prior Authorization Group** 

GEFITINIB, IRESSA **Drug Names** 

PA Indication Indicator

All FDA-approved Indications, Some Medically-accepted Indications

**IRESSA** 

Off-label Uses

Sensitizing epidermal growth factor receptor (EGFR) mutation-positive recurrent

non-small cell lung cancer (NSCLC).

**Exclusion Criteria** 

**Required Medical Information** 

For NSCLC: 1) disease must be metastatic, advanced, or recurrent and 2) patient must

have a sensitizing EGFR mutation.

Age Restrictions

**Coverage Duration** 

**Prescriber Restrictions** 

Plan Year

Other Criteria

Updated 07/01/2024 56 Prior Authorization GroupISTURISADrug NamesISTURISA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria - Required Medical Information - Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with an endocrinologist

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group IVERMECTIN TAB
Drug Names IVERMECTIN

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Ascariasis, Cutaneous larva migrans, Mansonelliasis, Scabies, Gnathostomiasis,

Pediculosis

Exclusion Criteria

**Required Medical Information** The requested drug is not being prescribed for the prevention or treatment of

coronavirus disease 2019 (COVID-19).

Age Restrictions --

Coverage Duration 1 month

Other Criteria -

Prior Authorization Group IWILFIN Drug Names IWILFIN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group JAKAFI
Drug Names JAKAFI

PA Indication Indicator
All FDA-approved Indications, Some Medically-accepted Indications
Off-label Uses
Lower-risk myelofibrosis, accelerated phase myelofibrosis, blast phase

myelofibrosis/acute myeloid leukemia, acute lymphoblastic leukemia (ALL), chronic myelomonocytic leukemia (CMML)-2, myelodysplastic syndrome/myeloproliferative neoplasm (MDS/MPN) with neutrophilia, essential thrombocythemia, and myeloid, lymphoid or mixed lineage neoplasms with eosinophilia and JAK2 rearrangement

Exclusion Criteria
Required Medical Information

For polycythemia vera: patient had an inadequate response or intolerance to interferon therapy or hydroxyurea. For acute lymphoblastic leukemia: patient has a cytokine receptor-like factor 2 (CRLF2) mutation or a mutation associated with activation of the Janus kinase/signal transducers and activators of transcription (JAK/STAT) pathway. For CMML-2: the requested drug is used in combination with a hypomethylating agent.

For myelodysplastic syndrome/myeloproliferative neoplasm (MDS/MPN) with neutrophilia: the requested drug is used as a single agent or in combination with a hypomethylating agent. For essential thrombocythemia: patient had an inadequate response or loss of response to hydroxyurea, interferon therapy, or anagrelide. For myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia and JAK2

rearrangement: the disease is in chronic or blast phase.

Age Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group JOENJA
Drug Names JOENJA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For activated phosphoinositide 3-kinase delta syndrome (APDS): the diagnosis was

confirmed by genetic testing demonstrating variant in either PIK3CD or PIK3R1.

Age Restrictions 12 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupJUXTAPIDDrug NamesJUXTAPID

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses
Exclusion Criteria
Required Medical Information

Required Medical Information

For initiation of therapy to treat homozygous familial hypercholesterolemia (HoFH), patient (pt) must meet ALL of the following: A) Diagnosis of HoFH confirmed by one of the following: 1) Genetic testing to confirm two mutant alleles at low-density lipoprotein receptor (LDLR), apolipoprotein B (ApoB), proprotein convertase subtilisin/kexin type 9 (PCSK9), or low-density lipoprotein receptor adaptor protein 1 (LDLRAP1) gene locus OR 2) History of an untreated low-density lipoprotein-cholesterol (LDL-C) of greater than 500 mg/dL or treated LDL-C greater than 300 mg/dL and either of the following: a) Presence of cutaneous or tendinous xanthomas before the age of 10 years, or b) An untreated LDL-C level of greater than or equal to 190 mg/dL in both parents, which is consistent with heterozygous familial hypercholesterolemia (HeFH), AND B) Prior to initiation of treatment, the pt is currently receiving treatment with a high-intensity statin at a maximally tolerated dose or at the maximum dose approved by the Food and Drug Administration (FDA) unless the pt is statin intolerant or has a contraindication to statin therapy, AND C) Prior to initiation of treatment with the requested drug, the pt is currently receiving treatment with a PCSK9-directed therapy at a maximally tolerated dose or at the maximum dose approved by the FDA unless the patient has experienced an intolerance or has a contraindication to all PCSK9-directed therapies, AND D) Prior to initiation of treatment, pt is/was experiencing an inadequate response to lipid-lowering therapy as indicated by a treated LDL-C greater than 100 mg/dL (or greater than 70 mg/dL with clinical atherosclerotic cardiovascular disease), AND E) The pt will continue to receive concomitant lipid lowering therapy. For renewal of therapy to treat HoFH: A) Pt meets all initial criteria, AND B) Has responded to therapy as demonstrated by a reduction in LDL-C from baseline, AND C) Is receiving concomitant lipid lowering therapy.

Age Restrictions
Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria

**Prior Authorization Group KALYDECO Drug Names KALYDECO** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

For cystic fibrosis (CF): The requested medication will not be used in combination with **Required Medical Information** 

other medications containing ivacaftor.

Age Restrictions **Prescriber Restrictions** 

**Coverage Duration** Plan Year

Other Criteria

**Prior Authorization Group** 

**Drug Names** PA Indication Indicator

Off-label Uses

**KANJINTI KANJINTI** 

All FDA-approved Indications, Some Medically-accepted Indications

Neoadiuvant treatment for human epidermal growth factor receptor 2 (HER2)-positive breast cancer, recurrent or advanced unresectable HER2-positive breast cancer, leptomeningeal metastases from HER2-positive breast cancer, brain metastases from HER2-positive breast cancer, HER2-positive esophageal and esophagogastric junction adenocarcinoma, HER2-positive advanced, recurrent, or metastatic uterine serous carcinoma, HER2-amplified and RAS and BRAF wild-type colorectal cancer (including appendiceal adenocarcinoma), HER2-positive recurrent salivary gland tumor, HER2-positive unresectable or metastatic hepatobiliary carcinoma (gallbladder cancer.

intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma), HER2 overexpression positive locally advanced, unresectable, or recurrent gastric

adenocarcinoma.

**Exclusion Criteria** 

Required Medical Information

All indications: the patient had an intolerable adverse event to Trazimera and that adverse event was NOT attributed to the active ingredient as described in the prescribing information. For colorectal cancer (including appendiceal adenocarcinoma): 1) the disease is HER2-amplified and RAS and BRAF wild-type and 2) the requested drug is used in combination with pertuzumab, tucatinib or lapatinib and 3) the patient has not had previous treatment with a HER2 inhibitor. For hepatobiliary carcinoma: 1) the disease is HER2-positive AND 2) the requested drug is used in combination with

pertuzumab.

Age Restrictions **Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Updated 07/01/2024 60 Prior Authorization Group KESIMPTA Drug Names KESIMPTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** KEVEYIS **Drug Names** KEVEYIS

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For primary HYPOkalemic periodic paralysis: 1) The diagnosis was supported by

genetic test results, OR 2) Patient has a family history of primary hypokalemic periodic

paralysis, OR 3) Patient's attacks are associated with hypokalemia AND both

Andersen-Tawil syndrome and thyrotoxic periodic paralysis have been ruled out. For primary HYPERkalemic periodic paralysis: 1) The diagnosis was supported by genetic

test results, OR 2) Patient has a family history of primary hyperkalemic periodic paralysis, OR 3) Patient's attacks are associated with hyperkalemia AND

Andersen-Tawil syndrome has been ruled out. For continuation of therapy for primary HYPOkalemic and primary HYPERkalemic periodic paralysis: Patient is demonstrating a response to therapy with the requested drug as demonstrated by a decrease in the

number or severity of attacks.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Initial: 2 months. Continuation: 12 months

Other Criteria -

Prior Authorization Group KEVZARA

Drug Names KEVZARA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For moderately to severely active rheumatoid arthritis (new starts only): 1) patient has

had an inadequate response, intolerance or contraindication to methotrexate (MTX) OR

2) patient has had an inadequate response or intolerance to a prior biologic

disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD. For polymyalgia rheumatica (PMR) (new starts only): 1) The patient has experienced an inadequate treatment response to corticosteroids OR 2) The patient has experienced a

disease flare while attempting to taper corticosteroids.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group KINERET Drug Names KINERET

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Systemic juvenile idiopathic arthritis, adult-onset Still's disease, multicentric Castleman's disease, Schnitzler syndrome, and Erdheim-Chester disease.

Exclusion Criteria -

Off-label Uses

**Required Medical Information** For moderately to severely active rheumatoid arthritis (new starts only): The patient has

experienced an inadequate treatment response, intolerance, or has a contraindication to two of the following products: adalimumab-aacf, Enbrel (etanercept), Humira

(adalimumab), Idacio (adalimumab-aacf), Kevzara (sarilumab), Rinvoq (upadacitinib), Xeljanz (tofacitinib)/Xeljanz XR (tofacitinib-extended release). For active systemic juvenile idiopathic arthritis (new starts only): patient must meet any of the following criteria: 1) Inadequate response to at least one nonsteroidal anti-inflammatory drug (NSAID), corticosteroid, methotrexate or leflunomide, 2) Inadequate response or intolerance to a prior biologic DMARD, OR 3) Physician global assessment score

greater than or equal to 5.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** KISQALI

**Drug Names** KISQALI, KISQALI FEMARA 200 DOSE, KISQALI FEMARA 400 DOSE, KISQALI

FEMARA 600 DOSE

PA Indication Indicator

Off-label Uses

All FDA-approved Indications, Some Medically-accepted Indications

Recurrent hormone receptor-positive, human epidermal growth factor receptor 2 (HER2)-negative breast cancer, in combination with an aromatase inhibitor, or

fulvestrant.

Exclusion Criteria -

**Required Medical Information** 

Age Restrictions -

Prescriber Restrictions - Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** KLISYRI **Drug Names** KLISYRI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** The patient has experienced an inadequate treatment response, intolerance, or has a

contraindication to ONE of the following: A) imiquimod 5 percent cream, B) fluorouracil

cream or solution.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** KORLYM

Drug NamesKORLYM, MIFEPRISTONEPA Indication IndicatorAll FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with an endocrinologist

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupKOSELUGODrug NamesKOSELUGO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses BRAF fusion or BRAF V600E activating mutation-positive recurrent or progressive

pilocytic astrocytoma

Exclusion Criteria -

Required Medical Information -

Age Restrictions For neurofibromatosis type 1: 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group KRAZATI
Drug Names KRAZATI

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** LAPATINIB

**Drug Names** LAPATINIB DITOSYLATE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Brain metastases from human epidermal growth factor receptor 2 (HER2)-positive breast cancer, recurrent HER2-positive breast cancer, recurrent epidermal growth factor receptor (EGFR)-positive chordoma, HER2-amplified and RAS and BRAF

wild-type colorectal cancer (including appendiceal adenocarcinoma).

Exclusion Criteria -

**Required Medical Information** For breast cancer, the patient meets all the following: a) the disease is recurrent,

advanced, or metastatic (including brain metastases), b) the disease is human

epidermal growth factor receptor 2 (HER2)-positive, c) the requested drug will be used in combination with any of the following: 1) aromatase inhibitor, 2) capecitabine, OR 3) trastuzumab. For colorectal cancer: 1) requested drug will be used in combination with trastuzumab and 2) patient has not had previous treatment with a HER2 inhibitor.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group Drug Names

LENVIMA

LENVIMA 10 MG DAILY DOSE, LENVIMA 12MG DAILY DOSE, LENVIMA 14 MG DAILY DOSE, LENVIMA 18 MG DAILY DOSE, LENVIMA 20 MG DAILY DOSE, LENVIMA 24 MG DAILY DOSE, LENVIMA 4 MG DAILY DOSE, LENVIMA 8 MG DAILY DOSE

PA Indication Indicator
Off-label Uses
Exclusion Criteria
Required Medical Information

All FDA-approved Indications, Some Medically-accepted Indications

Medullary thyroid carcinoma, recurrent endometrial carcinoma, thymic carcinoma

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For differentiated thyroid cancer (follicular, papillary, or Hurthle cell): disease is not amenable to radioactive iodine therapy and unresectable, locally recurrent, persistent, or metastatic. For hepatocellular carcinoma: disease is unresectable or inoperable, local, metastatic or with extensive liver tumor burden. For renal cell carcinoma, the disease is advanced, relapsed, or stage IV. For endometrial carcinoma, the patient meets ALL of the following: 1) The disease is advanced, recurrent, or metastatic, 2) The requested drug will be used in combination with pembrolizumab, 3) The patient experienced disease progression following prior systemic therapy, AND 4) The patient is not a candidate for curative surgery or radiation.

Age Restrictions
Prescriber Restrictions
Coverage Duration

Plan Year

Other Criteria -

**Prior Authorization Group** 

Drug Names

PA Indication Indicator

Off-label Uses

LEUPROLIDE

LEUPROLIDE ACETATE

All FDA-approved Indications, Some Medically-accepted Indications

Use in combination with growth hormone for children with growth failure and advancing puberty, recurrent androgen receptor positive salivary gland tumors, central precocious

puberty.

**Exclusion Criteria** 

**Required Medical Information** 

For central precocious puberty (CPP): Patients not currently receiving therapy must meet all of the following criteria: 1) Diagnosis of CPP was confirmed by a pubertal response to a gonadotropin releasing hormone (GnRH) agonist test OR a pubertal level

versus chronological age supports the diagnosis of CPP, AND 3) The onset of secondary sexual characteristics occurred prior to 8 years of age for female patients

of a third generation luteinizing hormone (LH) assay, AND 2) Assessment of bone age

OR prior to 9 years of age for male patients.

Age Restrictions CPP: Patient must be less than 12 years old if female and less than 13 years old if

male

Prescriber Restrictions

Coverage Duration

Plan Year

Other Criteria

**Prior Authorization Group** 

**Drug Names** 

LIDOCAINE PATCHES LIDOCAINE. LIDOCAN

PA Indication Indicator

Off-label Uses

All FDA-approved Indications, Some Medically-accepted Indications

Pain associated with diabetic neuropathy, pain associated with cancer-related

neuropathy (including treatment-related neuropathy [e.g., neuropathy associated with

radiation treatment or chemotherapy]).

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

LIQREV **LIQREV** 

**Prior Authorization Group Drug Names** 

PA Indication Indicator

All FDA-approved Indications

Off-label Uses

**Exclusion Criteria** 

**Required Medical Information** 

For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group

1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood

units.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

Updated 07/01/2024 66 Prior Authorization Group LIVMARLI Drug Names LIVMARLI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For treatment of cholestatic pruritis in a patient with Alagille syndrome (ALGS)

(continuation): the patient has experienced benefit from therapy (for example, improvement in pruritis). For treatment of cholestatic pruritis in a patient with

Progressive Familial Intrahepatic Cholestasis (PFIC), (initial): 1) diagnosis of PFIC has been confirmed by genetic testing, AND 2) the patient does not have PFIC type 2 with ABCB11 variants resulting in non-functional or complete absence of bile salt export pump (BSEP) protein, AND 3) the patient does not have any other concomitant liver disease, AND 4) the patient has not received a liver transplant. For treatment of cholestatic pruritis in a patient with PFIC, (continuation): the patient has experienced

benefit from therapy (for example, improvement in pruritis).

Age RestrictionsFor ALGS: 3 months of age or older, For PFIC: 5 years of age or olderPrescriber RestrictionsPrescribed by or in consultation with a hepatologist or gastroenterologist

Coverage Duration Initial: 6 months, Continuation: Plan Year

Other Criteria -

**Prior Authorization Group Drug Names**LIVTENCITY
LIVTENCITY

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -Required Medical Information -

**Age Restrictions** 12 years of age or older

**Prescriber Restrictions** Prescribed by or in consultation with an infectious disease specialist, transplant

specialist, hematologist, or oncologist.

Coverage Duration 3 months

Other Criteria -

Prior Authorization GroupLONSURFDrug NamesLONSURF

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For colorectal cancer (including appendiceal adenocarcinoma): The disease is

advanced or metastatic. For gastric or gastroesophageal junction adenocarcinoma, all of the following criteria must be met: 1) The disease is unresectable locally advanced, recurrent, or metastatic, and 2) The patient has been previously treated with at least

two prior lines of chemotherapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group LORBRENA
Drug Names LORBRENA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Anaplastic lymphoma kinase (ALK)-positive recurrent non-small cell lung cancer

(NSCLC). Repressor of silencing (ROS)-1 rearrangement-positive recurrent, advanced,

or metastatic NSCLC following progression on crizotinib, entrectinib, or ceritinib. Symptomatic or relapsed/refractory ALK-positive Erdheim-Chester Disease.

Inflammatory myofibroblastic tumor (IMT) with ALK translocation.

Exclusion Criteria -

**Required Medical Information** For recurrent, advanced, or metastatic NSCLC: Patient has ALK-positive disease.

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupLUMAKRASDrug NamesLUMAKRAS

PA Indication Indicator
All FDA-approved Indications, Some Medically-accepted Indications
Off-label Uses
Recurrent KRAS G12C-positive non-small cell lung cancer (NSCLC)

Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupLUMIZYMEDrug NamesLUMIZYME

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For Pompe disease: Diagnosis was confirmed by an enzyme assay demonstrating a

deficiency of acid alpha-glucosidase (GAA) enzyme activity or by genetic testing.

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupLUPKYNISDrug NamesLUPKYNIS

PA Indication Indicator All FDA-approved Indications

Off-label Uses -

**Exclusion Criteria**Use in combination with cyclophosphamide

**Required Medical Information** For lupus nephritis: 1) patient is currently receiving background immunosuppressive

therapy (e.g., mycophenolate mofetil, corticosteroids) for lupus nephritis, OR 2) patient has an intolerance or has a contraindication to background immunosuppressive therapy regimen for lupus nephritis. For lupus nephritis continuation: patient is receiving benefit from therapy and the benefit of continuing therapy outweighs the risk of worsening

nephrotoxicity.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** LUPRON-PROSTATE CA

**Drug Names** LUPRON DEPOT (1-MONTH), LUPRON DEPOT (3-MONTH), LUPRON DEPOT

(4-MONTH), LUPRON DEPOT (6-MONTH)

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Malignant sex cord-stromal tumors

Exclusion Criteria -

Required Medical Information -

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group LYNPARZA
Drug Names LYNPARZA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Recurrent HER2-negative, BRCA 1/2-germline mutated breast cancer, recurrent or metastatic HER2-positive, BRCA 1/2-germline mutated breast cancer, uterine

leiomyosarcoma.

Exclusion Criteria

**Required Medical Information** For recurrent or metastatic breast cancer: the disease is BRCA 1/2-germline mutated.

For prostate cancer: 1) The patient has a BRCA mutation and the requested drug will be used in combination with abiraterone and either prednisone or prednisolone OR 2) The patient has progressed on prior treatment with an androgen receptor-directed therapy. For epithelial ovarian, fallopian tube, or primary peritoneal cancer: The requested drug is used for maintenance therapy for stage II-IV or recurrent disease who are in complete or partial response to chemotherapy. For uterine leiomyosarcoma: 1) the patient has had at least one prior therapy AND 2) the patient has BRCA-altered

disease.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** MAVYRET **Drug Names** MAVYRET

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

**Exclusion Criteria** Decompensated cirrhosis/moderate or severe hepatic impairment (Child Turcotte Pugh

[CTP] class B or C).

**Required Medical Information** For hepatitis C virus (HCV): Infection confirmed by presence of HCV RNA in the serum

prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [CTP class B or C]), presence or absence of human immunodeficiency virus (HIV) coinfection, presence or absence of resistance-associated substitutions where applicable, transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current American Association for the Study of Liver Diseases and Infectious Diseases Society of America (AASLD-IDSA) treatment guidelines.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Criteria will be applied consistent with current AASLD-IDSA guidance

Other Criteria -

**Prior Authorization Group MEKINIST Drug Names MEKINIST** 

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Langerhans cell histiocytosis, Erdheim-Chester disease, Rosai-Dorfman disease.

**Exclusion Criteria** 

For melanoma: 1) The tumor is positive for a BRAF V600 activating mutation (e.g., **Required Medical Information** 

V600E or V600K). AND 2) The requested drug will be used as a single agent or in combination with dabrafenib. AND 3) The requested drug will be used for either of the following: a) unresectable, limited resectable, or metastatic disease, b) adjuvant systemic therapy. For central nervous system (CNS) cancer (i.e., glioma, oligodendroglioma, astrocytoma, glioblastoma), non-small cell lung cancer, solid tumors, and anaplastic thyroid cancer: 1) The tumor is positive for a BRAF V600E mutation, AND 2) The requested drug will be used in combination with dabrafenib. For uveal melanoma: The requested drug will be used as a single agent. For ovarian cancer, fallopian tube cancer, and primary peritoneal cancer: The requested drug will be used to treat persistent or recurrent disease. For gallbladder cancer, intrahepatic cholangiocarcinoma, and extrahepatic cholangiocarcinoma: 1) The tumor is positive for

a BRAF V600E mutation, AND 2) The disease is unresectable or metastatic, AND 3) The requested drug will be used in combination with dabrafenib. For papillary, follicular, and hurthle cell thyroid carcinoma: 1) The disease is positive for BRAF V600E

mutation, AND 2) The disease is not amenable to radioactive iodine (RAI) therapy.

AND 3) The requested drug will be used in combination with dabrafenib.

Age Restrictions **Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria

**Prior Authorization Group** MEKTOVI **Drug Names MEKTOVI** 

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Adjuvant systemic therapy for cutaneous melanoma, Langerhans Cell Histiocytosis

**Exclusion Criteria** 

**Required Medical Information** For melanoma: 1) The tumor is positive for BRAF V600 activating mutation (e.g., V600E or V600K), AND 2) The requested drug will be used in combination with encorafenib, AND 3) The requested drug will be used for either of the following: a)

unresectable, limited resectable, or metastatic disease, b) adjuvant systemic therapy.

**Age Restrictions Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria

Updated 07/01/2024 71 **Prior Authorization Group** MIGLUSTAT

Drug NamesMIGLUSTAT, YARGESAPA Indication IndicatorAll FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For type 1 Gaucher disease (GD1): Diagnosis was confirmed by an enzyme assay

demonstrating a deficiency of beta-glucocerebrosidase enzyme activity or by genetic

testing.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group MISCELLANEOUS INJECTABLES

**Drug Names** ABELCET, ACYCLOVIR SODIUM, AMPHOTERICIN B, AMPHOTERICIN B

**LIPOSOME** 

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Diagnosis of an FDA-approved indication not otherwise excluded from Part D.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group MODAFINIL
Drug Names MODAFINIL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For excessive sleepiness associated with narcolepsy: The diagnosis has been

confirmed by sleep lab evaluation. For excessive sleepiness associated with

obstructive sleep apnea (OSA): The diagnosis has been confirmed by

polysomnography.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupMOVANTIKDrug NamesMOVANTIK

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** Must have documented diagnosis of opioid-induced constipation caused by opioids

taken for chronic non-cancer pain, including patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage

escalation

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupMULPLETADrug NamesMULPLETA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For thrombocytopenia in patients with chronic liver disease: Untransfused platelet count

prior to a scheduled procedure is less than 50,000/mcL.

Age Restrictions 18 years of age or older

Prescriber Restrictions -

Coverage Duration 1 month

Other Criteria -

Prior Authorization Group
Drug Names

PA Indication Indicator
Off-label Uses

MVASI MVASI

All FDA-approved Indications, Some Medically-accepted Indications

Ampullary adenocarcinoma, breast cancer, central nervous system (CNS) cancers, malignant pleural mesothelioma, malignant peritoneal mesothelioma, pericardial mesothelioma, tunica vaginalis testis mesothelioma, soft tissue sarcomas, uterine neoplasms, endometrial carcinoma, vulvar cancers, small bowel adenocarcinoma, and ophthalmic-related disorders: diabetic macular edema, neovascular (wet) age-related macular degeneration including polypoidal choroidopathy and retinal angiomatous proliferation subtypes, macular edema following retinal vein occlusion, proliferative diabetic retinopathy, choroidal neovascularization, neovascular glaucoma and retinopathy of prematurity.

**Exclusion Criteria** 

**Required Medical Information** 

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For all indications except ophthalmic-related disorders: The patient had an intolerable adverse event to Zirabev and that adverse event was NOT attributed to the active ingredient as described in the prescribing information.

Age Restrictions
Prescriber Restrictions

Coverage Duration

Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization Group MYCAPSSA
Drug Names MYCAPSSA

PA Indication Indicator All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information

For acromegaly, initial: 1) Patient has a high pretreatment insulin-like growth factor-1 (IGF-1) level for age and/or gender based on the laboratory reference range, AND 2) Patient had an inadequate or partial response to surgery or radiotherapy OR there is a clinical reason for why the patient has not had surgery or radiotherapy. For acromegaly, continuation of therapy: Patient's IGF-1 level has decreased or normalized since

initiation of therapy.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupMYFEMBREEDrug NamesMYFEMBREE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information For heavy menstrual bleeding associated with uterine leiomyomas (fibroids) and

moderate to severe pain associated with endometriosis in a premenopausal patient: the patient has not already received greater than or equal to 24 months of treatment with

the requested drug.

Age Restrictions - Prescriber Restrictions -

Coverage Duration 12 months, max 24 months total

Other Criteria -

**Prior Authorization Group** MYTESI **Drug Names** MYTESI

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of non-infective diarrhea related to receiving

anti-retroviral therapy for HIV/AIDS. Must rule out any infectious types of diarrhea

before initiation.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NERLYNX
Drug Names NERLYNX

**PA Indication Indicator**All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent human epidermal growth factor receptor 2 (HER2)-positive breast cancer,

brain metastases from HER2-positive breast cancer.

Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group

**Drug Names** 

PA Indication Indicator

Off-label Uses

NEXAVAR

NEXAVAR, SORAFENIB TOSYLATE

All FDA-approved Indications, Some Medically-accepted Indications Acute myeloid leukemia, soft tissue sarcoma (angiosarcoma, desmoid

tumors/aggressive fibromatosis, and solitary fibrous tumor subtypes), gastrointestinal stromal tumor, medullary thyroid carcinoma, osteosarcoma, recurrent chordoma, epithelial ovarian cancer, fallopian tube cancer, primary peritoneal cancer, lymphoid,

myeloid, or mixed lineage neoplasms with eosinophilia

Exclusion Criteria

Required Medical Information

For acute myeloid leukemia: the disease is FMS-like tyrosine kinase 3-internal tandem duplication (FLT3-ITD) mutation-positive AND either of the following is met (1 OR 2): 1) the requested drug will be used as maintenance therapy after hematopoietic stem cell transplant, OR 2) the requested drug is used in combination with azacitidine or decitabine for low-intensity treatment induction or post-induction therapy AND either a) the patient has is 60 years of age or older or b) the disease is relapsed/refractory. For thyroid carcinoma: histology is follicular, papillary, Hurthle cell or medullary. For gastrointestinal stromal tumor (GIST): the patient meets either of the following: 1) the disease is unresectable, recurrent/progressive, or metastatic AND the patient has failed on an FDA-approved therapy (e.g., imatinib, sunitinib, regorafenib, ripretinib) OR 2) the requested drug is being used for palliation of symptoms if previously tolerated and effective. For renal cell carcinoma: the disease is advanced. For myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia: 1) the disease has a FLT3 rearrangement AND 2) the disease is in chronic or blast phase.

Age Restrictions -

**Prescriber Restrictions** 

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NEXLETOL

Drug Names NEXLETOL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupNEXLIZETDrug NamesNEXLIZET

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NGENLA
Drug Names NGENLA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

**Exclusion Criteria** Pediatric patients with closed epiphyses

**Required Medical Information** For pediatric growth hormone deficiency (GHD), initial: A) Patient (pt) has pre-treatment

(pre-tx) 1-year height (ht) velocity more than 2 standard deviations (SD) below mean OR a pre-tx ht more than 2 SD below mean and a 1-year ht velocity more than 1 SD below mean AND pt meets any of the following: 1) failed 2 pre-tx growth hormone (GH) stimulation tests (peak below 10 ng/mL), 2) pituitary/central nervous system (CNS) disorder (e.g., genetic defects, acquired structural abnormalities, congenital structural abnormalities) and pre-tx insulin-like growth factor-1 (IGF-1) more than 2 SD below

mean OR B) Pt was diagnosed with GHD as a neonate. For pediatric GHD,

continuation of therapy: Pt is experiencing improvement.

**Age Restrictions** 3 years of age or older

**Prescriber Restrictions** Prescribed by or in consultation with an endocrinologist

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NINLARO Drug Names NINLARO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Relapsed/refractory systemic light chain amyloidosis, Waldenstrom macroglobulinemia,

lymphoplasmacytic lymphoma

Exclusion Criteria -

Required Medical Information -

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** NITISINONE

Drug NamesNITISINONE, ORFADINPA Indication IndicatorAll FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For hereditary tyrosinemia type 1 (HT-1): Diagnosis of HT-1 is confirmed by one of the

following: 1) biochemical testing (e.g., detection of succinylacetone in urine) OR 2) DNA

testing (mutation analysis).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NORTHERA
Drug Names DROXIDOPA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information

For neurogenic orthostatic hypotension (nOH): Prior to initial therapy, patient has a persistent, consistent decrease in systolic blood pressure of at least 20 mmHg OR decrease in diastolic blood pressure of at least 10 mmHg within 3 minutes of standing or head-up tilt test. For continuation of therapy for nOH, patient must experience a sustained reduction in symptoms of nOH (i.e., decrease in dizziness, lightheadedness, or feeling faint). For both initial and continuation of therapy for nOH, the requested drug will be used for patients with neurogenic orthostatic hypotension associated with one of the following diagnoses: 1) primary autonomic failure due to Parkinson's disease, multiple system atrophy, or pure autonomic failure, OR 2) dopamine beta-hydroxylase deficiency, OR 3) non-diabetic autonomic neuropathy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration 3 months

Other Criteria -

Prior Authorization GroupNOXAFIL SUSPDrug NamesPOSACONAZOLE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** The requested drug will be used orally. For treatment of oropharyngeal candidiasis:

patient has experienced an inadequate treatment response, intolerance, or has a

contraindication to fluconazole.

**Age Restrictions** 13 years of age or older

Prescriber Restrictions -

Coverage Duration Oropharyngeal candidiasis: 1 month. All other indications: 6 months

Other Criteria -

Prior Authorization Group NUBEQA
Drug Names NUBEQA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** The requested drug will be used in combination with a gonadotropin-releasing hormone

(GnRH) analog or after bilateral orchiectomy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NI
Drug Names NI

PA Indication Indicator

Off-label Uses
Exclusion Criteria
Required Medical Information

NUCALA NUCALA

All FDA-approved Indications

For severe asthma, initial therapy: 1) Either a) Patient has baseline blood eosinophil count of at least 150 cells per microliter OR b) Patient is dependent on systemic corticosteroids, and 2) Patient has a history of severe asthma despite current treatment with both of the following medications: a) medium-to-high-dose inhaled corticosteroid and b) additional controller (i.e., long-acting beta2-agonist, long-acting muscarinic antagonist, leukotriene modifier, or sustained-release theophylline) unless patient has an intolerance or contraindication to such therapies. For severe asthma, continuation of therapy: Asthma control has improved on treatment with the requested drug, as demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations or a reduction in the daily maintenance oral corticosteroid dose. For eosinophilic granulomatosis with polyangiitis (EGPA), initial therapy: Patient has a history or the presence of an eosinophil count of more than 1000 cells per microliter or a blood eosinophil level of greater than 10 percent. For EGPA, continuation of therapy: Patient has a beneficial response to treatment with the requested drug, as

reduction in the daily oral corticosteroid dose, or 3) no active vasculitis. For hypereosinophilic syndrome (HES), initial therapy: 1) Patient has had HES for greater than or equal to 6 months, 2) Patient has HES without an identifiable non-hematologic secondary cause, 3) Patient does not have FIP1L1-PDGFRA kinase-positive HES, 4) Patient has a history or presence of a blood eosinophil count of at least 1000 cells per

demonstrated by any of the following: 1) a reduction in the frequency of relapses, 2) a

microliter, AND 5) Patient has been on a stable dose of at least one HES therapy (e.g., oral corticosteroid, immunosuppressive, and/or cytotoxic therapy). For HES, continuation of therapy: Patient has a beneficial response to treatment as

demonstrated by a reduction in HES flares.

Asthma: 6 years of age or older, EGPA and CRSwNP: 18 years of age or older, HES: 12 years of age or older

Prescriber Restrictions
Coverage Duration
Other Criteria

Age Restrictions

Plan Year

For chronic rhinosinusitis with nasal polyps (CRSwNP): 1) The requested drug is used as add-on maintenance treatment, AND 2) The patient has experienced inadequate treatment response to Xhance (fluticasone).

Prior Authorization GroupNUEDEXTADrug NamesNUEDEXTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria - Required Medical Information - Age Restrictions - Prescriber Restrictions - -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupNUPLAZIDDrug NamesNUPLAZID

PA Indication Indicator All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information For hallucinations and delusions associated with Parkinson's disease psychosis, the

diagnosis of Parkinson's disease must be made prior to the onset of psychotic

symptoms.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group NURTEC Drug Names NURTEC

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** Acute migraine treatment: The patient has experienced an inadequate treatment

response, intolerance, or the patient has a contraindication to one triptan 5-HT1 receptor agonist. Preventive treatment of migraine, initial: The patient meets either of the following: 1) The patient experienced an inadequate treatment response with a 4-week trial of any one of the following: Antiepileptic drugs (AEDs), Beta-adrenergic blocking agents, Antidepressants OR 2) The patient experienced an intolerance or has a contraindication that would prohibit a 4-week trial of any one of the following: Antiepileptic drugs (AEDs), Beta-adrenergic blocking agents, Antidepressants. Preventive treatment of migraine, continuation: The patient received at least 3 months of treatment with the requested drug, and the patient had a reduction in migraine days

per month from baseline.

Age Restrictions
Prescriber Restrictions

**Coverage Duration** Preventive treatment of migraine - initial: 3 months, All other indications: Plan Year

Other Criteria

Prior Authorization GroupOCALIVADrug NamesOCALIVA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For primary biliary cholangitis (PBC) without cirrhosis or with compensated cirrhosis

without evidence of portal hypertension: For initial therapy, 1) Diagnosis of PBC (previously known as primary biliary cirrhosis) is confirmed by at least two of the following: A) Biochemical evidence of cholestasis with elevation of alkaline phosphatase (ALP) level for at least 6 months duration, B) Presence of antimitochondrial antibodies (AMA) (titer greater than 1:40 by immunofluorescence or immunoenzymatic reactivity) or PBC-specific antinuclear antibodies ANA (eg, anti-gp210, anti-sp100), or C) Histologic evidence of PBC on liver biopsy (eg, non-suppurative inflammation and destruction of interlobular and septal bile ducts) and 2) Patient has an elevated serum ALP level prior to initiation of therapy with the requested drug and meets one of the following requirements: A) Inadequate response to at least 12 months of prior therapy with ursodeoxycholic acid (UDCA)/ursodiol and the patient will continue concomitant therapy with UDCA/ursodiol, or B) Intolerance to

UDCA/ursodiol. For continuation of therapy for PBC: patient achieved or maintained a

clinical benefit from Ocaliva therapy.

Age Restrictions - Prescriber Restrictions -

**Coverage Duration** Initial: 6 months. Continuation: Plan Year.

Other Criteria -

**Prior Authorization Group** OCTREOTIDE

**Drug Names** OCTREOTIDE ACETATE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Tumor control of thymomas and thymic carcinomas.

Exclusion Criteria -

**Required Medical Information** For acromegaly, initial: 1) Patient has a high pretreatment insulin-like growth factor-1

(IGF-1) level for age and/or gender based on the laboratory reference range, AND 2) Patient had an inadequate or partial response to surgery or radiotherapy OR there is a clinical reason for why the patient has not had surgery or radiotherapy. For acromegaly, continuation of therapy: Patient's IGF-1 level has decreased or normalized since initiation of therapy. For tumor control of thymomas and thymic carcinomas: The requested drug will be used for any of the following: 1) locally advanced or metastatic disease, 2) postoperatively following tumor resection.

Age Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prescriber Restrictions** 

Prior Authorization GroupODOMZODrug NamesODOMZO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** OFEV **Drug Names** OFEV

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For idiopathic pulmonary fibrosis (new starts only): 1) a high-resolution computed

tomography (HRCT) study of the chest or a lung biopsy reveals the usual interstitial pneumonia (UIP) pattern, OR 2) HRCT study of the chest reveals a result other than the UIP pattern (e.g., probable UIP, indeterminate for UIP) and the diagnosis is

supported either by a lung biopsy or by a multidisciplinary discussion between at least a radiologist and pulmonologist who are experienced in idiopathic pulmonary fibrosis if a

lung biopsy has not been conducted.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group
Drug Names
PA Indication Indicator

Off-label Uses

OGIVRI OGIVRI

All FDA-approved Indications, Some Medically-accepted Indications
Neoadjuvant treatment for human epidermal growth factor receptor 2 (HER2)-positive
breast cancer, recurrent or advanced unresectable HER2-positive breast cancer,
leptomeningeal metastases from HER2-positive breast cancer, brain metastases from
HER2-positive breast cancer, HER2-positive esophageal and esophagogastric junction
adenocarcinoma, HER2-positive advanced, recurrent, or metastatic uterine serous
carcinoma, HER2-amplified and RAS and BRAF wild-type colorectal cancer (including
appendiceal adenocarcinoma), HER2-positive recurrent salivary gland tumor,
HER2-positive unresectable or metastatic hepatobiliary carcinoma (gallbladder cancer,

intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma), HER2 overexpression positive locally advanced, unresectable, or recurrent gastric adenocarcinoma.

Exclusion Criteria
Required Medical Information

All indications: the patient had an intolerable adverse event to Trazimera and that adverse event was NOT attributed to the active ingredient as described in the prescribing information. For colorectal cancer (including appendiceal adenocarcinoma): 1) the disease is HER2-amplified and RAS and BRAF wild-type and 2) the requested drug is used in combination with pertuzumab, tucatinib or lapatinib and 3) the patient has not had previous treatment with a HER2 inhibitor. For hepatobiliary carcinoma: 1) the disease is HER2-positive AND 2) the requested drug is used in combination with

Age Restrictions Prescriber Restrictions Coverage Duration Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

pertuzumab.

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization GroupOGSIVEODrug NamesOGSIVEO

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupOJJAARADrug NamesOJJAARA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** 

**Drug Names** 

**OMNIPOD** 

All FDA-approved Indications

OMNIPOD 5 G6 INTRO KIT (G, OMNIPOD 5 G6 PODS (GEN 5), OMNIPOD CLASSIC PODS (GEN, OMNIPOD DASH INTRO KIT (G, OMNIPOD DASH PDM KIT (GEN,

OMNIPOD DASH PODS (GEN 4), OMNIPOD POD PALS

PA Indication Indicator

Off-label Uses -

Exclusion Criteria

**Required Medical Information** 

Omnipod GO, initial: 1) the patient has diabetes requiring insulin management AND 2)

the patient is currently self-testing glucose levels, the patient will be counseled on self-testing glucose levels, or the patient is using a continuous glucose monitor AND 3) the patient has experienced an inadequate treatment response or intolerance to long-acting basal insulin therapy. Omnipod, V-GO, initial: 1) The patient has diabetes requiring insulin management with multiple daily injections AND 2) The patient is self-testing glucose levels 4 or more times per day OR the patient is using a continuous glucose monitor AND 3) The patient has experienced any of the following with the current diabetes regimen: inadequate glycemic control, recurrent hypoglycemia, wide fluctuations in blood glucose, dawn phenomenon with persistent severe early morning

hyperglycemia, severe glycemic excursions.

Age Restrictions -

**Prescriber Restrictions** 

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ONUREG
Drug Names ONUREG

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupOPSUMITDrug NamesOPSUMIT

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group

1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) Pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood

units.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** 

**Drug Names** 

**PA Indication Indicator** 

Off-label Uses
Exclusion Criteria

Required Medical Information

ORAL-INTRANASAL FENTANYL

FENTANYL CITRATE, FENTANYL CITRATE ORAL TRA, LAZANDA, SUBSYS

All FDA-approved Indications

1) The requested drug is indicated for the treatment of breakthrough cancer-related pain only. The requested drug is being prescribed for the management of breakthrough

pain in a cancer patient with underlying cancer pain AND 2) The International Classification of Diseases (ICD) diagnosis code provided supports the cancer-related diagnosis. [Note: For drug coverage approval, ICD diagnosis code provided MUST support the cancer-related diagnosis.] AND 3) The patient is currently receiving, and will continue to receive, around-the-clock opioid therapy for underlying cancer pain AND 4) The requested drug is intended only for use in opioid tolerant patients. The patient can safely take the requested dose based on their current opioid use history. [Note: Patients considered opioid tolerant are those who are taking around-the-clock medicine consisting of at least 60 mg of oral morphine per day, at least 25 mcg per hour of transdermal fentanyl, at least 30 mg of oral oxycodone per day, at least 60 mg of oral hydrocodone per day, at least 25 mg of oral oxymorphone per day, or an equianalgesic dose of another opioid

medication daily for one week or longer.].

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**Prior Authorization Group** 

**Drug Names** 

PA Indication Indicator

Off-label Uses

**Exclusion Criteria** 

**ORALAIR** 

Plan Year

**ORALAIR** 

UNALAIN

All FDA-approved Indications

Severe, unstable or uncontrolled asthma. History of any severe systemic allergic

reaction or any severe local reaction to sublingual allergen immunotherapy. History of

eosinophilic esophagitis.

**Required Medical Information** 

Age Restrictions

-

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5 to 65 years of age

**Prescriber Restrictions** 

Prescribed by or in consultation with an allergist or immunologist

**Coverage Duration** 

Plan Year

Other Criteria

**Prior Authorization Group** ORENITRAM

**Drug Names** ORENITRAM, ORENITRAM TITRATION KIT M

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For pulmonary arterial hypertension (World Health Organization [WHO] Group 1): PAH

was confirmed by right heart catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment

pulmonary vascular resistance is greater than or equal to 3 Wood units.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupORGOVYXDrug NamesORGOVYX

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria - Required Medical Information - Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ORIAHNN
Drug Names ORIAHNN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information For heavy menstrual bleeding associated with uterine leiomyomas (fibroids) in a

premenopausal patient: the patient has not already received greater than or equal to 24

months of treatment with any elagolix-containing drug.

Age Restrictions -

Prescriber Restrictions

Coverage Duration 12 months, max 24 months total

Other Criteria -

Prior Authorization GroupORKAMBIDrug NamesORKAMBI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For cystic fibrosis (CF): The requested medication will not be used in combination with

other medications containing ivacaftor.

**Age Restrictions** 1 year of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupORLADEYODrug NamesORLADEYO

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the prevention of acute angioedema attacks due to hereditary angioedema (HAE):

The patient meets either of the following: 1) the patient has hereditary angioedema (HAE) with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing OR 2) the patient has hereditary angioedema with normal C1 inhibitor confirmed by laboratory testing and either of the following: a) patient tested positive for an F12, angiopoietin-1, plasminogen, kininogen-1 (KNG1), heparan sulfate-glucosamine 3-O-sulfotransferase 6 (HS3ST6), or myoferlin (MYOF) gene mutation OR b) patient has a family history of angioedema and the angioedema was refractory to a trial of high-dose antihistamine

therapy for at least one month.

Age Restrictions 12 years of age or older

Prescriber Restrictions Prescribed by or in consultation with an immunologist, allergist, or rheumatologist

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupOTEZLADrug NamesOTEZLA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupOXANDROLONEDrug NamesOXANDROLONE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Cachexia associated with AIDS (HIV wasting). To enhance growth in patients with

**Turners Syndrome** 

Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Turners Syndrome: Plan Year, All other diagnoses: 6 months

Other Criteria Coverage will be denied if request is for an indication excluded from Medicare Part D.

Prior Authorization Group OXERVATE
Drug Names OXERVATE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with an ophthalmologist or optometrist

Coverage Duration 8 weeks
Other Criteria -

Prior Authorization GroupOZEMPICDrug NamesOZEMPIC

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria The Prior Authorization only applies to patients whose claim is not submitted with an

ICD-10 code indicating a diagnosis of type 2 diabetes mellitus OR to patients who do not have a history of an antidiabetic drug (EXCLUDING glucagon-like peptide receptor agonists [GLP-1 RAs] and combination glucose-dependent insulinotropic polypeptide

[GIP] and GLP-1 RAs).

Prior Authorization GroupPALYNZIQDrug NamesPALYNZIQ

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupPANRETINDrug NamesPANRETIN

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Topical treatment of cutaneous lesions in patients with non-AIDS-related Kaposi

sarcoma

Exclusion Criteria -

Required Medical Information - Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** PAROXETINE SUSP

Drug NamesPAROXETINE HYDROCHLORIDEPA Indication IndicatorAll FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** Patient is unable to take solid oral dosage forms (e.g., difficulty swallowing tablets or

capsules).

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group
PEGASYS
PEGASYS

**PA Indication Indicator**All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Myeloproliferative neoplasm (essential thrombocythemia, polycythemia vera, symptomatic lower-risk myelofibrosis), systemic mastocytosis, adult T-cell

leukemia/lymphoma, mycosis fungoides/sezary syndrome, primary cutaneous CD30+ T-cell lymphoproliferative disorders, hairy cell leukemia, Erdheim-Chester disease.

initial treatment during pregnancy for chronic myeloid leukemia.

Exclusion Criteria

Required Medical Information For chronic hepatitis C: Hepatitis C virus (HCV) confirmed by presence of hepatitis C

virus HCV RNA in serum prior to starting treatment and the planned treatment regimen.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration HCV: 12-48wks. Criteria applied consistent w/current AASLD/IDSA guidance. HBV:

48wks. Other: Plan Yr

Other Criteria -

Prior Authorization GroupPEMAZYREDrug NamesPEMAZYRE

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** PHENYLBUTYRATE

**Drug Names** OLPRUVA, SODIUM PHENYLBUTYRATE

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For urea cycle disorders (UCD): Diagnosis of UCD was confirmed by enzymatic,

biochemical, or genetic testing.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** PIQRAY

**Drug Names** PIQRAY 200MG DAILY DOSE, PIQRAY 250MG DAILY DOSE, PIQRAY 300MG

**DAILY DOSE** 

PA Indication Indicator

Off-label Uses

All FDA-approved Indications, Some Medically-accepted Indications

Recurrent hormone receptor (HR)-positive, human epidermal growth factor receptor 2

(HER2)-negative, PIK3CA-mutated breast cancer in combination with fulvestrant.

Exclusion Criteria -

Required Medical Information

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group POMALYST

Drug Names POMALYST

PA Indication Indicator

Off-label Uses

All FDA-approved Indications, Some Medically-accepted Indications

Relapsed/refractory systemic light chain amyloidosis, primary central nervous system

(CNS) lymphoma, POEMS (polyneuropathy, organomegaly, endocrinopathy,

monoclonal protein, skin changes) syndrome.

Exclusion Criteria

**Required Medical Information** 

For multiple myeloma, patient has previously received at least two prior therapies for

multiple myeloma, including an immunomodulatory agent AND a proteasome inhibitor.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group PREVYMIS

Drug Names PREVYMIS

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** 

For prophylaxis of cytomegalovirus (CMV) infection or disease in hematopoietic stem

cell transplant (HSCT): 1) the patient is CMV-seropositive, AND 2) the patient is a recipient of an allogeneic HSCT. For prophylaxis of CMV disease in kidney transplant: 1) the patient is CMV-seronegative, AND 2) the patient is a high risk recipient of kidney

transplant.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration 7 months

Other Criteria

Prior Authorization Group Drug Names PA Indication Indicator

Off-label Uses

PROCRIT PROCRIT

All FDA-approved Indications, Some Medically-accepted Indications

Anemia due to myelodysplastic syndromes (MDS), anemia in rheumatoid arthritis (RA), anemia due to hepatitis C treatment (ribavirin in combination with either interferon alfa or peginterferon alfa)

Exclusion Criteria
Required Medical Information

Patients receiving chemotherapy with curative intent. Patients with myeloid cancer. Requirements regarding hemoglobin (Hgb) values exclude values due to a recent transfusion. For initial approval: 1) for all uses except anemia due to chemotherapy or myelodysplastic syndrome (MDS): patient has adequate iron stores (for example, a transferrin saturation [TSAT] greater than or equal to 20%), AND 2) for all uses except surgery: pretreatment (no erythropoietin treatment in previous month) Hgb is less than 10 g/dL, AND 3) for MDS: pretreatment serum erythropoietin level is 500 international units/L or less. For reauthorizations (patient received erythropoietin treatment in previous month) in all uses except surgery: 1) patient has received at least 12 weeks of erythropoietin therapy, AND 2) patient responded to erythropoietin therapy, AND 3) current Hgb is less than 12 g/dL, AND 4) for all uses except anemia due to chemotherapy or MDS: patient has adequate iron stores (for example, a transferrin saturation [TSAT] greater than or equal to 20%).

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

\_

16 weeks

Coverage includes use in anemia in patients whose religious beliefs forbid blood transfusions. Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual (e.g., used for treatment of anemia for a patient with chronic renal failure who is undergoing dialysis, or furnished from physician's supply incident to a physician service).

**Prior Authorization Group PROLIA PROLIA Drug Names** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria Required Medical Information** 

For postmenopausal osteoporosis, patient (pt) has ONE of the following: 1) history of fragility fracture, OR 2) pre-treatment (pre-tx) T-score of less than or equal to -2.5 or pre-tx T-score greater than -2.5 and less than -1 with a high pre-tx Fracture Risk

Assessment Tool (FRAX) fracture probability AND pt has ANY of the following: a) indicators for higher fracture risk (e.g., advanced age, frailty, glucocorticoid therapy, very low T-scores, or increased fall risk), b) pt has failed prior treatment with or is intolerant to a previous injectable osteoporosis therapy, or c) pt has had an oral bisphosphonate trial of at least 1-year duration or there is a clinical reason to avoid treatment with an oral bisphosphonate. For osteoporosis in men: pt has one of the following: 1) history of osteoporotic vertebral or hip fracture, OR 2) pre-tx T-score of less than or equal to -2.5 or pre-tx T-score greater than -2.5 and less than -1 with a high pre-tx FRAX fracture probability AND patient has ANY of the following: a) patient has failed prior treatment with or is intolerant to a previous injectable osteoporosis therapy, OR b) patient has had an oral bisphosphonate trial of at least 1-year duration or there is a clinical reason to avoid treatment with an oral bisphosphonate. For glucocorticoid-induced osteoporosis: 1) pt has had an oral bisphosphonate trial of at least 1-year duration unless pt has a contraindication or intolerance to an oral bisphosphonate, AND 2) pt has one of the following: a) history of fragility fracture, OR b) pre-tx T-score of less than or equal to -2.5, OR c) pre-tx T-score greater than -2.5 and less than -1 with a high pre-tx FRAX fracture probability. For breast cancer, pt is receiving adjuvant aromatase inhibitor therapy. For prostate cancer, pt is receiving androgen deprivation therapy (ADT).

Age Restrictions **Prescriber Restrictions Coverage Duration** Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual. Patient has high FRAX fracture probability if the 10 year probability is either greater than or equal to 20 percent for any major osteoporotic fracture or greater than or equal to 3 percent for hip fracture. If glucocorticoid treatment is greater than 7.5 mg (prednisone equivalent) per day, the estimated risk score generated with FRAX should be multiplied by 1.15 for major osteoporotic fracture and 1.2 for hip fracture.

Updated 07/01/2024 96 Prior Authorization Group PROMACTA

PROMACTA

PROMACTA

**PA Indication Indicator** All FDA-a<sub>l</sub>

Off-label Uses
Exclusion Criteria
Required Medical Information

PROMACTA

All FDA-approved Indications

For chronic or persistent immune thrombocytopenia (ITP): 1) For new starts: a) Patient (pt) has had an inadequate response or is intolerant to a prior therapy such as corticosteroids or immunoglobulins AND b) Untransfused platelet (plt) count at any point prior to the initiation of the requested medication is less than 30.000/mcL OR 30,000-50,000/mcL with symptomatic bleeding or risk factor(s) for bleeding (e.g., undergoing a medical or dental procedure where blood loss is anticipated. comorbidities such as peptic ulcer disease and hypertension, anticoagulation therapy, profession or lifestyle that predisposes pt to trauma) AND c) For chronic ITP only: pt has had an inadequate response or intolerance to Doptelet (avatrombopag). 2) For continuation of therapy, plt count response to the requested drug: a) Current plt count is less than or equal to 200,000/mcL, OR b) Current plt count is greater than 200,000/mcL to less than or equal to 400,000/mcL and dosing will be adjusted to a plt count sufficient to avoid clinically important bleeding. For thrombocytopenia associated with chronic hepatitis C: 1) For new starts: the requested drug is used for initiation and maintenance of interferon-based therapy. 2) For continuation of therapy: pt is receiving interferon-based therapy. For severe aplastic anemia (AA): 1) For new starts: a) Pt will use the requested drug with standard immunosuppressive therapy for first line treatment OR b) the pt had an insufficient response to immunosuppressive therapy. 2) For continuation of therapy: 1) Current plt count is 50,000-200,000/mcL, OR 2) Current

plt count is less than 50.000/mcL and pt has not received appropriately titrated therapy

transfusion-independent, OR 4) Current plt count is greater than 200,000/mcL to less than or equal to 400.000/mcL and dosing will be adjusted to achieve and maintain an

for at least 16 weeks, OR 3) Current plt count is less than 50,000/mcL and pt is

Age Restrictions
Prescriber Restrictions

**Coverage Duration** 

Other Criteria

HCV: 6mo, ITP/AA initial: 6mo, ITP reauth: Plan Year, AA reauth: APR-Plan Year,

IPR-16 wks

APR: adequate platelet response (greater than 50,000/mcL), IPR: inadequate platelet

response (less than 50,000/mcL).

appropriate target plt count.

**Prior Authorization Group** PYRUKYND

**Drug Names** PYRUKYND, PYRUKYND TAPER PACK

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For hemolytic anemia in a patient with pyruvate kinase (PK) deficiency: Diagnosis was

confirmed by an enzyme assay demonstrating deficiency of PK enzyme activity or by genetic testing. For hemolytic anemia in a patient with PK deficiency (continuation of therapy): Patient achieved or maintained a positive clinical response (e.g., improvement

in hemoglobin levels, reduction in blood transfusions).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Initial: 7 months, Continuation: Plan Year

Other Criteria -

Prior Authorization GroupQINLOCKDrug NamesQINLOCK

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent/progressive or unresectable gastrointestinal stromal tumor (GIST)

Exclusion Criteria -

**Required Medical Information** For unresectable, recurrent/progressive, advanced, or metastatic gastrointestinal

stromal tumor (GIST), the patient meets either of the following: 1) patient has received prior treatment with 3 or more kinase inhibitors, including imatinib OR 2) patient has experienced disease progression following treatment with avapritinib and dasatinib.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupQUININE SULFATEDrug NamesQUININE SULFATE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Babesiosis, uncomplicated Plasmodium vivax malaria.

Exclusion Criteria -

**Required Medical Information** For babesiosis: the requested drug is used in combination with clindamycin.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration 1 month

Other Criteria -

**Prior Authorization Group** RADICAVA

**Drug Names** RADICAVA, RADICAVA ORS STARTER KIT

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For amyotrophic lateral sclerosis (ALS): 1) Diagnosis is classified as definite or

probable ALS, AND 2) For new starts only: Patient has scores of at least 2 points on all 12 areas of the revised ALS Functional Rating Scale (ALSFRS-R). For continuation of

therapy for ALS: There is a clinical benefit from therapy.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group RAVICTI
Drug Names RAVICTI

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For urea cycle disorders (UCD): Diagnosis of UCD was confirmed by enzymatic,

biochemical or genetic testing.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupRECORLEVDrug NamesRECORLEV

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with an endocrinologist

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupRELISTOR INJDrug NamesRELISTOR

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the treatment of opioid-induced constipation in a patient with chronic non-cancer

pain, including chronic pain related to prior cancer or its treatment who does not require frequent (e.g., weekly) opioid dosage escalation: 1) the patient is unable to tolerate oral medications OR 2) the patient meets one of the following criteria A) experienced an inadequate treatment response or intolerance to an oral drug indicated for opioid-induced constipation in a patient with chronic non-cancer pain (e.g., Movantik) OR B) the patient has a contraindication that would prohibit a trial of an oral drug

indicated for opioid-induced constipation in a patient with chronic non-cancer pain (e.g.,

Movantik).

Age Restrictions - Prescriber Restrictions -

Coverage Duration 4 months

Other Criteria -

Prior Authorization Group RELISTOR TAB
Drug Names RELISTOR

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration 4 months

Other Criteria -

Prior Authorization Group Drug Names PA Indication Indicator Off-label Uses

Exclusion Criteria
Required Medical Information

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

**REMICADE** 

INFLIXIMAB, REMICADE

All FDA-approved Indications, Some Medically-accepted Indications
Behcet's syndrome, hidradenitis suppurativa, juvenile idiopathic arthritis, pyoderma
gangrenosum, sarcoidosis, Takayasu's arteritis, uveitis.

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For moderately to severely active rheumatoid arthritis (new starts only): 1) Pt meets ANY of the following: a) requested drug will be used in combination with methotrexate (MTX) or leflunomide OR b) intolerance or CI to MTX AND leflunomide, AND 2) Pt meets ANY of the following: a) inadequate response, intolerance or CI to MTX OR b) inadequate response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD. For active ankylosing spondylitis (new starts only): an inadequate treatment response or intolerance to a non-steroidal anti-inflammatory drug (NSAID) OR contraindication that would prohibit a trial of NSAIDs. For moderate to severe plague psoriasis (new starts only): 1) At least 3% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin, intertriginous areas) are affected at time of diagnosis, AND 2) Pt meets ANY of the following: a) pt has experienced inadequate response or intolerance to either phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with MTX, cyclosporine, or acitretin, OR b) pharmacologic treatment with MTX, cyclosporine, or acitretin is contraindicated, OR c) pt has severe psoriasis that warrants a biologic as first-line therapy (i.e., at least 10% of BSA or crucial body areas [e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas] are affected).

Plan Year

For hidradenitis suppurativa (new starts only): Pt has severe, refractory disease. For uveitis (new starts only): Inadequate response or intolerance or has a CI to a trial of immunosuppressive therapy for uveitis. For FDA-approved indications and off-label uses that overlap: The patient had an intolerable adverse event to Renflexis and that adverse event was NOT attributed to the active ingredient as described in the prescribing information.

**Prior Authorization Group Drug Names** 

PA Indication Indicator

Off-label Uses

**RENFLEXIS** All FDA-approved Indications, Some Medically-accepted Indications

**RENFLEXIS** 

Behcet's syndrome, hidradenitis suppurativa, juvenile idiopathic arthritis, pyoderma

gangrenosum, sarcoidosis, Takayasu's arteritis, uveitis

**Exclusion Criteria Required Medical Information** 

For moderately to severely active rheumatoid arthritis (new starts only): 1) Pt meets

ANY of the following: a) requested drug will be used in combination with methotrexate (MTX) or leflunomide OR b) intolerance or CI to MTX AND leflunomide, AND 2) pt meets ANY of the following: a) inadequate response, intolerance or CI to MTX OR b) inadequate response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD. For active ankylosing spondylitis (new starts only): an inadequate treatment response or intolerance to a non-steroidal anti-inflammatory drug (NSAID) OR contraindication that would prohibit a trial of NSAIDs. For moderate to severe plague psoriasis (new starts only): 1) At least 3% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin, intertriginous areas) are affected at time of diagnosis, AND 2) Pt meets ANY of the following: a) pt has experienced inadequate response or intolerance to either phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with MTX, cyclosporine, or acitretin, OR b) pharmacologic treatment with MTX, cyclosporine, or acitretin is contraindicated, OR c) pt has severe psoriasis that warrants a biologic as first-line therapy (i.e., at least 10% of BSA or crucial body areas [e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas] are affected).

Age Restrictions **Prescriber Restrictions Coverage Duration** Other Criteria

Plan Year

For hidradenitis suppurativa (new starts only): pt has severe, refractory disease. For uveitis (new starts only): Inadequate response or intolerance or has a CI to a trial of

immunosuppressive therapy for uveitis.

**Prior Authorization Group** 

**Drug Names** 

REPATHA

REPATHA. REPATHA PUSHTRONEX SYSTEM. REPATHA SURECLICK

All FDA-approved Indications

Off-label Uses

**Exclusion Criteria** 

PA Indication Indicator

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

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Prior Authorization Group Drug Names PA Indication Indicator

Off-label Uses

RETACRIT RETACRIT

All FDA-approved Indications, Some Medically-accepted Indications

Anemia due to myelodysplastic syndromes (MDS), anemia in rheumatoid arthritis (RA), anemia due to hepatitis C treatment (ribavirin in combination with either interferon alfa or peginterferon alfa)

Exclusion Criteria
Required Medical Information

Patients receiving chemotherapy with curative intent. Patients with myeloid cancer. Requirements regarding hemoglobin (Hgb) values exclude values due to a recent transfusion. For initial approval: 1) for all uses except anemia due to chemotherapy or myelodysplastic syndrome (MDS): patient has adequate iron stores (for example, a transferrin saturation [TSAT] greater than or equal to 20%), AND 2) for all uses except surgery: pretreatment (no erythropoietin treatment in previous month) Hgb is less than 10 g/dL, AND 3) for MDS: pretreatment serum erythropoietin level is 500 international units/L or less. For reauthorizations (patient received erythropoietin treatment in previous month) in all uses except surgery: 1) patient has received at least 12 weeks of erythropoietin therapy, AND 2) patient responded to erythropoietin therapy, AND 3) current Hgb is less than 12 g/dL, AND 4) for all uses except anemia due to chemotherapy or MDS: patient has adequate iron stores (for example, a transferrin saturation [TSAT] greater than or equal to 20%).

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

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16 weeks

Coverage includes use in anemia in patients whose religious beliefs forbid blood transfusions. Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual (e.g., used for treatment of anemia for a patient with chronic renal failure who is undergoing dialysis, or furnished from physician's supply incident to a physician service).

Prior Authorization GroupRETEVMODrug NamesRETEVMOPA Indication IndicatorAll FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Recurrent rearranged during transfection (RET)-rearrangement positive non-small cell lung cancer, Langerhans Cell Histiocytosis with a RET gene fusion, symptomatic or relapsed/refractory Erdheim-Chester Disease with a RET gene fusion, symptomatic or relapsed/refractory Rosai-Dorfman Disease with a RET gene fusion, RET-fusion positive recurrent or persistent thyroid carcinoma (papillary carcinoma, follicular carcinoma, and Hurthle cell carcinoma), RET-fusion positive anaplastic thyroid carcinoma.

Exclusion Criteria

**Required Medical Information** 

For non-small cell lung cancer, patient must meet all of the following: 1) The disease is recurrent, advanced or metastatic, and 2) Tumor is RET fusion-positive or RET rearrangement-positive.

Medullary thyroid cancer and thyroid cancer: 12 years of age or older.

Age Restrictions
Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria

**Prior Authorization Group** 

**Drug Names** 

PA Indication Indicator

Off-label Uses

**REVLIMID** 

LENALIDOMIDE. REVLIMID

All FDA-approved Indications, Some Medically-accepted Indications

Systemic light chain amyloidosis, classical Hodgkin lymphoma, myelodysplastic syndrome without the 5g deletion cytogenetic abnormality, myelofibrosis-associated anemia, POEMS (polyneuropathy, organomegaly, endocrinopathy, monoclonal protein, skin changes) syndrome, myeloproliferative neoplasms, Kaposi Sarcoma, Langerhans

cell histiocytosis, peripheral T-Cell lymphomas not otherwise specified.

angioimmunoblastic T-cell lymphoma (AITL), enteropathy-associated T-cell lymphoma,

monomorphic epitheliotropic intestinal T-cell lymphoma, nodal peripheral T-cell lymphoma, adult T-cell leukemia/lymphoma, hepatosplenic T-cell lymphoma, primary central nervous system (CNS) lymphoma, chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL), acquired immunodeficiency syndrome (AIDS)-related B-cell lymphoma, monomorphic post-transplant lymphoproliferative disorder, diffuse large B-cell lymphoma, multicentric Castleman's disease, high-grade B-cell

lymphoma.

**Exclusion Criteria** 

**Required Medical Information** 

For myelodysplastic syndrome (MDS): patient has lower risk MDS with symptomatic

lymphomas, histologic transformation of indolent lymphoma to diffuse large B-cell

anemia per the Revised International Prognostic Scoring System (IPSS-R),

International Prognostic Scoring System (IPSS), or World Health organization (WHO)

classification-based Prognostic Scoring System (WPSS).

Age Restrictions

**Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria

**Prior Authorization Group** REZUROCK **Drug Names REZUROCK** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions 12 years of age or older

**Prescriber Restrictions** 

**Coverage Duration** Plan Year

Other Criteria

Updated 07/01/2024 105 **Prior Authorization Group RINVOQ RINVOQ Drug Names** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria Required Medical Information** 

For moderately to severely active rheumatoid arthritis (new starts only): patient has experienced an inadequate treatment response, intolerance or has a contraindication to at least one tumor necrosis factor (TNF) inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]), For active psoriatic arthritis (new starts only): patient has experienced an inadequate treatment response. intolerance or has a contraindication to at least one TNF inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]). For moderately to severely active ulcerative colitis (new starts only): patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one TNF inhibitor (e.g., adalimumab-aacf, Humira [adalimumab], Idacio [adalimumab-aacf]). For moderately to severely active Crohn's disease (new starts only): patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one TNF inhibitor (e.g., adalimumab-aacf, Humira [adalimumab], Idacio [adalimumab-aacf]). For atopic dermatitis (new starts only): 1) patient has refractory, moderate to severe disease, AND 2) patient has had an inadequate response to treatment with other systemic drug products, including biologics, or use of these therapies are inadvisable. For atopic dermatitis (continuation of therapy): the patient achieved or maintained positive clinical response. For active ankylosing spondylitis (new starts only): patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one TNF inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]). For non-radiographic axial spondyloarthritis (new starts only): patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one TNF inhibitor.

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Atopic dermatitis: 12 years of age or older

Atopic dermatitis (initial): 4 months, All others: Plan Year

Other Criteria

Updated 07/01/2024 106 Prior Authorization GroupRIVFLOZADrug NamesRIVFLOZA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For primary hyperoxaluria type 1 (PH1): diagnosis has been confirmed by a molecular

genetic test showing a mutation in the alanine:glyoxylate aminotransferase (AGXT)

gene or liver enzyme analysis demonstrating absent or significantly reduced

alanine:glyoxylate aminotransferase (AGT) activity. For PH1 (continuation): the patient has experienced decreased or normalized levels of urinary oxalate since initiating

therapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ROZLYTREK
Drug Names ROZLYTREK

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Recurrent ROS1-positive non-small cell lung cancer (NSCLC), Non-metastatic neurotrophic tyrosine receptor kinase (NTRK) gene fusion-positive solid tumors,

first-line treatment of NTRK gene fusion-positive solid tumors.

Exclusion Criteria -

**Required Medical Information** For all neurotrophic tyrosine receptor kinase (NTRK) gene fusion-positive solid tumors,

the disease is without a known acquired resistance mutation. For ROS1-positive

non-small cell lung cancer, the patient has recurrent, advanced, or metastatic disease.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupRUBRACADrug NamesRUBRACA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Uterine leiomyosarcoma, pancreatic adenocarcinoma, advanced (stage II-IV) epithelial

ovarian, fallopian tube, or primary peritoneal cancer

Exclusion Criteria
Required Medical Information

For metastatic castration-resistant prostate cancer with a deleterious breast cancer susceptibility gene (BRCA) mutation (germline and/or somatic): 1) patient has been treated with androgen receptor-directed therapy, AND 2) patient has been treated with a taxane-based chemotherapy or the patient is not fit for chemotherapy, AND 3) the requested drug will be used in combination with a gonadotropin-releasing hormone (GnRH) analog or after bilateral orchiectomy. For maintenance treatment of BRCA mutated epithelial ovarian, fallopian tube, primary peritoneal cancer: 1) the patient has advanced (stage II-IV) disease and is in complete or partial response to primary therapy, OR 2) the patient has recurrent disease and is in complete or partial response to platinum-based chemotherapy. For uterine leiomyosarcoma: 1) the requested drug is used as second-line therapy, AND 2) the patient has BRCA-altered disease. For pancreatic adenocarcinoma: 1) the patient has metastatic disease, AND 2) the patient has somatic or germline BRCA or PALB-2 mutations.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupRYBELSUSDrug NamesRYBELSUS

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria

The Prior Authorization only applies to patients whose claim is not submitted with an ICD-10 code indicating a diagnosis of type 2 diabetes mellitus OR to patients who do not have a history of an antidiabetic drug (EXCLUDING glucagon-like peptide receptor agonists [GLP-1 RAs] and combination glucose-dependent insulinotropic polypeptide

[GIP] and GLP-1 RAs).

Prior Authorization Group RYDAPT Drug Names RYDAPT

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Relapsed or refractory acute myeloid leukemia (AML), myeloid, lymphoid, or mixed

lineage neoplasms with eosinophilia and FGFR1 or FLT3 rearrangements, post-induction therapy for AML, re-induction in residual disease for AML

Exclusion Criteria -

**Required Medical Information** For acute myeloid leukemia (AML): AML is FMS-like tyrosine kinase 3 (FLT3)

mutation-positive. For myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia and Fibroblast growth factor receptor type 1 (FGFR1) or FLT3 rearrangements: the

disease is in chronic or blast phase.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** SAPROPTERIN

**Drug Names** JAVYGTOR, SAPROPTERIN DIHYDROCHLORI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information

For phenylketonuria (PKU): For patients who have not yet received a therapeutic trial of the requested drug, the patient's pretreatment (including before dietary management) phenylalanine level is greater than 6 mg/dL (360 micromol/L). For patients who completed a therapeutic trial of the requested drug, the patient must have experienced improvement (e.g., reduction in blood phenylalanine levels, improvement in

neuropsychiatric symptoms).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Initial: 2 months, All others: Plan Year

Other Criteria -

**Prior Authorization Group** SAVELLA

**Drug Names** SAVELLA, SAVELLA TITRATION PACK

PA Indication Indicator All FDA-approved Indications

Off-label Uses -

Exclusion Criteria -

**Required Medical Information** The patient has experienced an inadequate treatment response, intolerance, or has a

contraindication to duloxetine or pregabalin.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSCEMBLIXDrug NamesSCEMBLIX

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For chronic myeloid leukemia (CML) in the chronic phase: 1) the diagnosis was

confirmed by detection of the Philadelphia chromosome or BCR-ABL gene AND 2) the patient meets either of the following: A) the patient has previously been treated with 2 or more tyrosine kinase inhibitors (TKIs) AND at least one of those was imatinib or

dasatinib, OR B) the patient is positive for the T315I mutation.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSEROSTIMDrug NamesSEROSTIM

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the treatment of human immunodeficiency virus (HIV) patients with wasting or

cachexia: The requested medication is used in combination with antiretroviral therapy. Patient has had a suboptimal response to at least one other therapy for wasting or cachexia (e.g., megestrol, dronabinol, cyproheptadine, or testosterone therapy if hypogonadal) or patient has a contraindication or intolerance to alternative therapies. For continuation of therapy, patient must have demonstrated a response to therapy with the requested medication (i.e., body mass index [BMI] has increased or stabilized).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration 12 weeks

Other Criteria -

Prior Authorization Group SIGNIFOR Drug Names SIGNIFOR

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions -

Prescriber Restrictions Prescribed by or in consultation with an endocrinologist

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** SILDENAFIL

**Drug Names** SILDENAFIL CITRATE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group

1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1) Pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) Pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) If the request is for an adult, pretreatment pulmonary vascular resistance is

greater than or equal to 3 Wood units.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSIRTURODrug NamesSIRTURO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions -

**Prescriber Restrictions** Prescribed by or in consultation with an infectious disease specialist.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group SKYCLARYS
Drug Names SKYCLARYS

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For Friedreich's ataxia (FRDA): 1) The patient has a confirmed genetic mutation in the

frataxin (FXN) gene, AND 2) The patient is exhibiting clinical manifestations of the disease (e.g., muscle weakness, decline in coordination, frequent falling). For FRDA continuation of therapy: The patient has experienced a beneficial response to therapy

(e.g., slowing of clinical decline).

Age Restrictions 16 years of age or older

**Prescriber Restrictions** Prescribed by or in consultation with a physician who specializes in Friedreich's ataxia

or a neurologist

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** SKYRIZI

Drug Names SKYRIZI, SKYRIZI PEN

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For moderate to severe plaque psoriasis (new starts only): at least 3% of body surface

area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin,

intertriginous areas) are affected at the time of diagnosis.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSOHONOSDrug NamesSOHONOS

PA Indication Indicator All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For reduction in volume of new heterotopic ossification in fibrodysplasia ossificans

progressiva (FOP): The patient has a confirmed genetic mutation in the activin A

receptor type I (ACVR1) gene.

Age Restrictions 8 years of age or older if female and 10 years of age or older if male

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSOMAVERTDrug NamesSOMAVERT

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For acromegaly, initial: 1) Patient has a high pretreatment insulin-like growth factor-1 (IGF-1) level for age and/or gender based on the laboratory reference range, AND 2)

Patient had an inadequate or partial response to surgery or radiotherapy OR there is a clinical reason for why the patient has not had surgery or radiotherapy. For acromegaly,

continuation of therapy: Patient's IGF-1 level has decreased or normalized since

initiation of therapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** 

Drug Names

PA Indication Indicator

Off-label Uses

SPRYCEL SPRYCEL

All FDA-approved Indications, Some Medically-accepted Indications

Gastrointestinal stromal tumor (GIST), metastatic chondrosarcoma, recurrent chordoma, T-cell acute lymphoblastic leukemia (ALL), and Philadelphia (Ph)-like B-ALL, myeloid and/or lymphoid neoplasms with eosinophilia and ABL1 rearrangement

in the chronic phase or blast phase

used for palliation of symptoms.

**Exclusion Criteria** 

**Required Medical Information** 

For chronic myeloid leukemia (CML), including patients who have received a hematopoietic stem cell transplant: 1) Diagnosis was confirmed by detection of the Philadelphia (Ph) chromosome or BCR-ABL gene, and 2) If patient experienced resistance to an alternative tyrosine kinase inhibitor, patient is negative for all of the following mutations: T315I/A, F317L/V/I/C, and V299L. For acute lymphoblastic leukemia (ALL), the patient has a diagnosis of one of the following: 1) Philadelphia chromosome positive ALL, including patients who have received a hematopoietic stem cell transplant: diagnosis that has been confirmed by detection of the Ph chromosome or BCR-ABL gene, and if patient experienced resistance to an alternative tyrosine kinase inhibitor, patient is negative for all of the following mutations: T315I/A, F317L/V/I/C, and V299L, OR 2) Ph-like B-ALL with ABL-class kinase fusion, OR 3) relapsed or refractory T-cell ALL with ABL-class kinase fusion. For GIST, 1) the patient meets all of the following: A) the disease is unresectable, recurrent/progressive, or metastatic, B) the patient has received prior therapy with imatinib or avapritinib AND C)

patients is positive for PDGFRA exon 18 mutations, OR 2) the requested drug is being

Age Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prescriber Restrictions** 

Prior Authorization Group STELARA
Drug Names STELARA

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For moderate to severe plaque psoriasis (new starts): At least 3% of body surface area

(BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin,

intertriginous areas) are affected at the time of diagnosis.

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group STIVARGA
Drug Names STIVARGA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Progressive gastrointestinal stromal tumors (GIST), osteosarcoma, glioblastoma,

angiosarcoma, retroperitoneal/intra-abdominal soft tissue sarcoma,

rhabdomyosarcoma, soft tissue sarcomas of the extremities, body wall, head and neck.

Exclusion Criteria -

**Required Medical Information** For gastrointestinal stromal tumors: The disease is progressive, locally advanced,

unresectable, or metastatic. For colorectal cancer: The disease is advanced or

metastatic.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSUNOSIDrug NamesSUNOSI

PA Indication Indicator All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For excessive daytime sleepiness associated with narcolepsy, initial request: 1) The

diagnosis has been confirmed by sleep lab evaluation, AND 2) The patient has experienced an inadequate treatment response or intolerance to at least one central nervous system (CNS) wakefulness promoting drug (e.g., armodafinil, modafinil), OR has a contraindication that would prohibit a trial of central nervous system (CNS) wakefulness promoting drugs (e.g., armodafinil, modafinil). For excessive daytime sleepiness associated with obstructive sleep apnea (OSA), initial request: 1) The diagnosis has been confirmed by polysomnography, AND 2) The patient has

experienced an inadequate treatment response or intolerance to at least one central nervous system (CNS) wakefulness promoting drug (e.g., armodafinil, modafinil), OR has a contraindication that would prohibit a trial of central nervous system (CNS) wakefulness promoting drugs (e.g., armodafinil, modafinil). If the request is for a

continuation of therapy, then the patient experienced a decrease in daytime sleepiness with narcolepsy or a decrease in daytime sleepiness with obstructive sleep apnea

(OSA).

Age Restrictions

Prescriber Restrictions

Coverage Duration
Other Criteria

Prescribed by or in consultation with a sleep disorder specialist or neurologist

Plan Year

**Prior Authorization Group** SUTENT

**Drug Names** SUNITINIB MALATE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Thyroid carcinoma (follicular medullary papillary and Hurthle cell)

Thyroid carcinoma (follicular, medullary, papillary, and Hurthle cell), soft tissue sarcoma (angiosarcoma, solitary fibrous tumor, and alveolar soft part sarcoma subtypes), recurrent chordoma, thymic carcinoma, lymphoid, myeloid, or mixed lineage neoplasms with eosinophilia, pheochromocytoma, paraganglioma, gastrointestinal stromal tumor (GIST) (unresectable, recurrent/progressive, or metastatic disease after progression on approved therapies, unresectable succinate dehydrogenase (SDH)-deficient GISTs and

use for palliation of symptoms if previously tolerated and effective).

Exclusion Criteria
Required Medical Information

For renal cell carcinoma (RCC): the patient meets either of the following: 1) the disease is relapsed, advanced, or stage IV OR 2) the requested drug is being used as adjuvant treatment for patients that are at high risk of recurrent RCC following nephrectomy. For gastrointestinal stromal tumor (GIST): the patient meets one of the following: 1) the requested drug will be used after disease progression on or intolerance to imatinib, 2) the disease is unresectable, recurrent/progressive, or metastatic AND the patient has failed on an FDA-approved therapy (e.g., imatinib, sunitinib, regorafenib, ripretinib), 3) the requested drug will be used for unresectable succinate dehydrogenase (SDH)-deficient GIST, OR 4) the requested drug will be used for the palliation of symptoms if previously tolerated and effective. For myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia: 1) the disease has a FLT3 rearrangement AND 2) the disease is in chronic or blast phase.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group SYMDEKO
Drug Names SYMDEKO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

Exclusion Criteria -

**Required Medical Information** For cystic fibrosis: The requested medication will not be used in combination with other

medications containing ivacaftor.

Age Restrictions 6 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupSYMPAZANDrug NamesSYMPAZAN

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Seizures associated with Dravet syndrome

Exclusion Criteria -

Required Medical Information

Age Restrictions Seizures associated with Lennox-Gastaut syndrome (LGS): 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupTABRECTADrug NamesTABRECTA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent non-small cell lung cancer (NSCLC).

Exclusion Criteria -

**Required Medical Information** For recurrent, advanced, or metastatic NSCLC: Tumor is positive for

mesenchymal-epithelial transition (MET) exon 14 skipping mutation.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** TADALAFIL (BPH)

**Drug Names** TADALAFIL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of Benign prostatic hyperplasia.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupTADALAFIL (PAH)Drug NamesALYQ, TADALAFIL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information For pulmonary arterial hypertension (PAH) (World Health Organization [WHO] Group

1): PAH was confirmed by right heart catheterization. For PAH new starts only: 1)
Pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2)
Pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg,
AND 3) Pretreatment pulmonary vascular resistance is greater than or equal to 3 Wood

units.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group Drug Names PA Indication Indicator Off-label Uses TAFINLAR TAFINLAR

All FDA-approved Indications, Some Medically-accepted Indications
Thyroid carcinoma (papillary carcinoma, follicular carcinoma, and Hurthle cell
carcinoma), central nervous system (CNS) cancer (i.e., oligodendroglioma,
astrocytoma, glioblastoma), gallbladder cancer, extrahepatic cholangiocarcinoma,
intrahepatic cholangiocarcinoma, Langerhans cell histiocytosis, Erdheim-Chester
disease, ovarian cancer, fallopian tube cancer, and primary peritoneal cancer.

Exclusion Criteria
Required Medical Information

For central nervous system (CNS) cancer (i.e., glioma, oligodendroglioma, astrocytoma, glioblastoma): 1) The tumor is positive for a BRAF V600E mutation AND 2) The requested drug will be used in combination with trametinib. For melanoma: 1) The tumor is positive for a BRAF V600 activating mutation (e.g., V600E or V600K), AND 2) The requested drug will be used as a single agent or in combination with trametinib, AND 3) The requested drug will be used for either of the following: a) unresectable, limited resectable, or metastatic disease, b) adjuvant systemic therapy. For non-small cell lung cancer: 1) The tumor is positive for a BRAF V600E mutation, AND 2) The requested drug will be used as a single agent or in combination with trametinib. For papillary, follicular, and Hurthle cell thyroid carcinoma: 1) The tumor is BRAF-positive, AND 2) The disease is not amenable to radioactive iodine (RAI) therapy. For Langerhans Cell Histiocytosis and Erdheim-Chester Disease: The disease is positive for a BRAF V600E mutation. For gallbladder cancer, extrahepatic cholangiocarcinoma, and intrahepatic cholangiocarcinoma: 1) The disease is positive for a BRAF V600E mutation, AND 2) The disease is unresectable or metastatic, AND 3) The requested drug will be used in combination with trametinib. For solid tumors: 1) The tumor is positive for a BRAF V600E mutation, AND 2) The requested drug will be used in combination with trametinib. For ovarian cancer, fallopian tube cancer, and primary peritoneal cancer: 1) The disease is positive for BRAF V600E mutation, AND 2) The disease is persistent or recurrent, AND 3) The requested drug will be used in combination with trametinib.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

-

Plan Year

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Prior Authorization Group TAGRISSO
Drug Names TAGRISSO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Sensitizing epidermal growth factor receptor (EGFR) mutation-positive recurrent non-small cell lung cancer (NSCLC), brain metastases from sensitizing EGFR mutation-positive NSCLC, leptomeningeal metastases from EGFR mutation-positive

NSCLC.

Exclusion Criteria -

**Required Medical Information** For NSCLC, the requested drug is used in any of the following settings: 1) The patient

meets both of the following: a) patient has metastatic, advanced, or recurrent NSCLC (including brain and/or leptomeningeal metastases from NSCLC) and b) patient has a sensitizing EGFR mutation OR 2) The patient meets both of the following: a) request is for adjuvant treatment of NSCLC following tumor resection and b) patient has EGFR

mutation-positive disease.

\*\*Age Restrictions\*\*
-

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupTAKHZYRODrug NamesTAKHZYRO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the prevention of acute angioedema attacks due to hereditary angioedema (HAE):

The patient meets either of the following: 1) the patient has hereditary angioedema (HAE) with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing OR 2) the patient has hereditary angioedema with normal C1 inhibitor confirmed by laboratory testing and either of the following: a) patient tested positive for an F12, angiopoietin-1, plasminogen, kininogen-1 (KNG1), heparan sulfate-glucosamine 3-O-sulfotransferase 6 (HS3ST6), or myoferlin (MYOF) gene mutation OR b) patient has a family history of angioedema and the angioedema was refractory to a trial of high-dose antihistamine

therapy for at least one month.

Age Restrictions 2 years of age or older

Prescriber Restrictions Prescribed by or in consultation with an immunologist, allergist, or rheumatologist

Coverage Duration Plan Year
Other Criteria -

Prior Authorization Group TALTZ
Drug Names TALTZ

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** 

For moderate to severe plague psoriasis (new starts only): 1) at least 3% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin, intertriginous areas) are affected at the time of diagnosis AND 2) the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to one of the following products: adalimumab-aacf, Enbrel (etanercept), Humira (adalimumab), Idacio (adalimumab-aacf), Otezla (apremilast), Skyrizi (risankizumab-rzaa), Stelara (ustekinumab), Tremfya (guselkumab). For active ankylosing spondylitis (new starts only): the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to one of the following products: adalimumab-aacf, Enbrel (etanercept), Humira (adalimumab), Idacio (adalimumab-aacf), Rinvog (upadacitinib), Xelianz (tofacitinib)/Xelianz XR (tofacitinib) extended-release). For active psoriatic arthritis (PsA) (new starts only): the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to one of the following products: adalimumab-aacf, Enbrel (etanercept), Humira (adalimumab), Idacio (adalimumab-aacf), Otezla (apremilast), Rinvog (upadacitinib), Skyrizi (risankizumab-rzaa), Stelara (ustekinumab), Tremfya (guselkumab), Xeljanz (tofacitinib)/Xeljanz XR (tofacitinib extended-release). For active non-radiographic axial spondyloarthritis (new starts only): patient meets any of the following: 1) patient has experienced an inadequate treatment response to a non-steroidal anti-inflammatory drug (NSAID) OR 2) patient has experienced an intolerance or has a contraindication to NSAIDs.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TALZENNA
Drug Names TALZENNA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent germline breast cancer susceptibility gene (BRCA)-mutated breast cancer

Exclusion Criteria -

Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** 

**Drug Names** 

PA Indication Indicator

Off-label Uses

TARGRETIN TOPICAL

**BEXAROTENE** 

All FDA-approved Indications, Some Medically-accepted Indications

Stage 2 or higher mycosis fungoides (MF)/Sezary syndrome (SS), chronic or

smoldering adult T-cell leukemia/lymphoma (ATLL), primary cutaneous marginal zone

lymphoma, primary cutaneous follicle center lymphoma

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Carrage Described

Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

PA Indication Indicator

Off-label Uses

TASIGNA TASIGNA

All FDA-approved Indications, Some Medically-accepted Indications

Philadelphia chromosome positive acute lymphoblastic leukemia (Ph+ ALL), gastrointestinal stromal tumor (GIST), myeloid and/or lymphoid neoplasms with eosinophilia and ABL1 rearrangement in the chronic phase or blast phase, pigmented

villonodular synovitis/tenosynovial giant cell tumor

**Exclusion Criteria** 

**Required Medical Information** 

For chronic myeloid leukemia (CML), including patients newly diagnosed with CML and patients who have received a hematopoietic stem cell transplant, 1) Diagnosis was

confirmed by detection of the Philadelphia chromosome or BCR-ABL gene, 2) patient has experienced resistance or intolerance to imatinib or dasatinib, AND 3) If patient experienced resistance to an alternative tyrosine kinase inhibitor for CML, patient is

negative for T315I, Y253H, E255K/V, and F359V/C/I mutations. For acute

lymphoblastic leukemia (ALL), including patients who have received a hematopoietic stem cell transplant: 1) Diagnosis was confirmed by detection of the Philadelphia chromosome or BCR-ABL gene, AND 2) if the patient has experienced resistance to an alternative tyrosine kinase inhibitor for ALL, patient is negative for T315I, Y253H,

E255K/V, F359V/C/I and G250E. For gastrointestinal stromal tumor (GIST), the

patients meets either of the following: 1) the disease is unresectable,

recurrent/progressive, or metastatic AND the disease has progressed on at least 2 approved therapies (e.g. imatinib, sunitinib, dasatinib, regorafenib, ripretinib) OR 2) the

requested drug is being prescribed for palliation of symptoms.

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Plan Year

Other Criteria

**Prior Authorization Group TAVNEOS TAVNEOS Drug Names** 

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** For continuation of treatment for severe anti-neutrophil cytoplasmic autoantibody

(ANCA)-associated vasculitis: the patient has experienced benefit from therapy.

Age Restrictions **Prescriber Restrictions** 

**Coverage Duration** Plan Year

Other Criteria

**Prior Authorization Group TAZAROTENE** 

**Drug Names** TAZAROTENE. TAZORAC PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** For plaque psoriasis, the patient meets the following criteria: 1) the patient has less

> than or equal to 20 percent of affected body surface area (BSA), AND 2) the patient experienced an inadequate treatment response or intolerance to at least one topical

corticosteroid OR has a contraindication that would prohibit a trial of topical

corticosteroids.

Age Restrictions **Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria

**Prior Authorization Group TAZVERIK Drug Names TAZVERIK** 

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions Epithelioid sarcoma: 16 years of age or older, Follicular lymphoma: 18 years of age or

older

**Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria

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**Prior Authorization Group** TECFIDERA

**Drug Names** DIMETHYL FUMARATE, DIMETHYL FUMARATE STARTER

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prescriber Restrictions** 

Prior Authorization Group TEGSEDI Drug Names TEGSEDI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For polyneuropathy of hereditary transthyretin-mediated amyloidosis initial therapy,

patient is positive for a mutation of the TTR gene and exhibits clinical manifestation of

disease. For polyneuropathy of hereditary transthyretin-mediated amyloidosis

continuation, patient demonstrates a beneficial response to therapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TEPMETKO
Drug Names TEPMETKO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent non-small cell lung cancer (NSCLC).

Exclusion Criteria -

**Required Medical Information** For recurrent, advanced, or metastatic NSCLC: Tumor is positive for

mesenchymal-epithelial transition (MET) exon 14 skipping mutation.

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** 

Drug Names

PA Indication Indicator

PA Indication Indicator

Off-label Uses
Exclusion Criteria
Required Medical Information

TERIPARATIDE TERIPARATIDE

All FDA-approved Indications

For postmenopausal osteoporosis: patient has ONE of the following: 1) history of fragility fracture, OR 2) pre-treatment T-score of less than or equal to -2.5 or pre-treatment T-score greater than -2.5 and less than -1 with a high pre-treatment Fracture Risk Assessment Tool (FRAX) fracture probability AND patient has ANY of the following: a) indicators for higher fracture risk (e.g., advanced age, frailty, glucocorticoid therapy, very low T-scores, or increased fall risk). OR b) patient has failed prior treatment with or is intolerant to a previous injectable osteoporosis therapy OR c) patient has had an oral bisphosphonate trial of at least 1-year duration or there is a clinical reason to avoid treatment with an oral bisphosphonate. For primary or hypogonadal osteoporosis in men: patient has ONE of the following: 1) history of osteoporotic vertebral or hip fracture, OR 2) pre-treatment T-score of less than or equal to -2.5, or pre-treatment T-score greater than -2.5 and less than -1 with a high pre-treatment FRAX fracture probability AND patient has ANY of the following: a) patient has failed prior treatment with or is intolerant to a previous injectable osteoporosis therapy, OR b) patient has had an oral bisphosphonate trial of at least 1-year duration or there is a clinical reason to avoid treatment with an oral bisphosphonate.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Initial: 24 months, Continuation: Plan Year

For glucocorticoid-induced osteoporosis: Patient has had an oral bisphosphonate trial of at least 1-year duration unless patient has a contraindication or intolerance to an oral bisphosphonate, AND patient meets ANY of the following: 1) patient has a history of fragility fracture, OR 2) a pre-treatment T-score of less than or equal to -2.5, OR 3) pre-treatment T-score greater than -2.5 and less than -1 with a high pre-treatment FRAX fracture probability. Continuation of therapy: If the patient has received greater than or equal to 24 months of therapy with any parathyroid hormone analog: 1) The patient remains at or has returned to having a high risk for fracture, AND 2) The benefit of therapy with this prescribed medication outweighs the potential risks for this patient.

**Prior Authorization Group** 

**Drug Names** 

PA Indication Indicator

Off-label Uses

**Exclusion Criteria Required Medical Information**  **TESTOSTERONE CYPIONATE INJ** 

DEPO-TESTOSTERONE. TESTOSTERONE CYPIONATE

All FDA-approved Indications, Some Medically-accepted Indications

Gender Dysphoria

For primary hypogonadism or hypogonadotropic hypogonadism, initial therapy: The patient has at least two confirmed low morning serum total testosterone concentrations based on the reference laboratory range or current practice guidelines [Note: Safety and efficacy of testosterone products in patients with "age-related hypogonadism" (also referred to as "late-onset hypogonadism") have not been established.]. For primary hypogonadism or hypogonadotropic hypogonadism, continuation of therapy: The patient had a confirmed low morning serum total testosterone concentration based on the reference laboratory range or current practice guidelines before starting testosterone therapy [Note: Safety and efficacy of testosterone products in patients with "age-related hypogonadism" (also referred to as "late-onset hypogonadism") have not been established.]. For gender dysphoria: The patient is able to make an informed decision to engage in hormone therapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**PA Indication Indicator** 

Off-label Uses

**Exclusion Criteria** 

Required Medical Information

TESTOSTERONE ENANTHATE INJ TESTOSTERONE ENANTHATE

All FDA-approved Indications, Some Medically-accepted Indications

Gender Dysphoria

For primary hypogonadism or hypogonadotropic hypogonadism, initial therapy: The patient has at least two confirmed low morning serum total testosterone concentrations based on the reference laboratory range or current practice guidelines [Note: Safety and efficacy of testosterone products in patients with "age-related hypogonadism" (also referred to as "late-onset hypogonadism") have not been established.]. For primary hypogonadism or hypogonadotropic hypogonadism, continuation of therapy: The patient had a confirmed low morning serum total testosterone concentration based on the reference laboratory range or current practice guidelines before starting testosterone therapy [Note: Safety and efficacy of testosterone products in patients with "age-related hypogonadism" (also referred to as "late-onset hypogonadism") have not been established.]. For gender dysphoria: The patient is able to make an informed decision to engage in hormone therapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

Updated 07/01/2024 125 Prior Authorization Group TETRABENAZINE
Drug Names TETRABENAZINE

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Tic disorders, tardive dyskinesia, hemiballismus, chorea not associated with

Huntington's disease.

Exclusion Criteria

**Required Medical Information** For treatment of tardive dyskinesia and treatment of chorea associated with

Huntington's disease: The patient has experienced an inadequate treatment response

or intolerable adverse event to deutetrabenazine.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group THALOMID
Drug Names THALOMID

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Myelofibrosis-associated anemia, AIDS-related aphthous stomatitis, Kaposi sarcoma,

chronic graft-versus-host disease, Crohn's disease, multicentric Castleman's disease,

Rosai-Dorfman disease, Langerhans cell histiocytosis

Exclusion Criteria

Required Medical Information -

Age Restrictions

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TIBSOVO
Drug Names TIBSOVO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Conventional (grades 1-3) or dedifferentiated chondrosarcoma. Newly-diagnosed acute

myeloid leukemia (AML) if 60-74 years of age and without comorbidities.

Exclusion Criteria
Required Medical Information

Patient has disease with a susceptible isocitrate dehydrogenase-1 (IDH1) mutation. For acute myeloid leukemia (AML): 1) patient has newly-diagnosed AML and meets one of the following: a) 75 years of age or older, b) patient has comorbidities that preclude use of intensive induction chemotherapy, or c) patient is 60 years of age or older and declines intensive induction chemotherapy, OR 2) patient is 60 years of age or older and the requested drug will be used as post-induction therapy following response to induction therapy with the requested drug, OR 3) patient has relapsed or refractory AML. For locally advanced, unresectable, or metastatic cholangiocarcinoma: the requested drug will be used as subsequent treatment for progression on or after systemic treatment.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TOPICAL TESTOSTERONES

**Drug Names** TESTOSTERONE, TESTOSTERONE PUMP

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Gender Dysphoria

Exclusion Criteria -

For primary hypogonadism or hypogonadotropic hypogonadism, initial therapy: The patient has at least two confirmed low morning serum total testosterone concentrations based on the reference laboratory range or current practice guidelines [Note: Safety and efficacy of testosterone products in patients with "age-related hypogonadism" (also referred to as "late-onset hypogonadism") have not been established.]. For primary hypogonadism or hypogonadotropic hypogonadism, continuation of therapy: The patient had a confirmed low morning serum total testosterone concentration based on the reference laboratory range or current practice guidelines before starting testosterone therapy [Note: Safety and efficacy of testosterone products in patients with

"age-related hypogonadism" (also referred to as "late-onset hypogonadism") have not been established.]. For gender dysphoria: The patient is able to make an informed

decision to engage in hormone therapy.

Age Restrictions
Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupTOPICAL TRETINOINDrug NamesAVITA, TRETINOIN

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group

Drug Names

PA Indication Indicator

Off-label Uses

TRAZIMERA TRAZIMERA

All FDA-approved Indications, Some Medically-accepted Indications

Neoadjuvant treatment for human epidermal growth factor receptor 2 (HER2)-positive breast cancer, recurrent or advanced unresectable HER2-positive breast cancer, leptomeningeal metastases from HER2-positive breast cancer, brain metastases from HER2-positive breast cancer, brain metastases from HER2-positive breast cancer, HER2-positive esophageal and esophagogastric junction adenocarcinoma, HER2-positive advanced, recurrent, or metastatic uterine serous carcinoma, HER2-amplified and RAS and BRAF wild-type colorectal cancer (including appendiceal adenocarcinoma), HER2-positive recurrent salivary gland tumor.

HER2-positive unresectable or metastatic hepatobiliary carcinoma (gallbladder cancer.

intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma), HER2 overexpression positive locally advanced, unresectable, or recurrent gastric

adenocarcinoma.

Exclusion Criteria

**Required Medical Information** 

For colorectal cancer (including appendiceal adenocarcinoma): 1) the disease is HER2-amplified and RAS and BRAF wild-type and 2) the requested drug is used in combination with pertuzumab, tucatinib or lapatinib and 3) the patient has not had previous treatment with a HER2 inhibitor. For hepatobiliary carcinoma: 1) the disease is HER2 positive and 2) the requested drug is used in combination with pertuzumab.

Age Restrictions Prescriber Restrictions -

**Coverage Duration** 

Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** TRELSTAR

**Drug Names** TRELSTAR MIXJECT

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Gender dysphoria, ovarian suppression in breast cancer

Exclusion Criteria -

**Required Medical Information** For gender dysphoria, patient meets either of the following (1 or 2): 1) the requested

drug is used to suppress puberty and the patient is at Tanner stage 2 or greater, OR 2) patient is undergoing gender transition, and the patient will receive the requested drug concomitantly with gender-affirming hormones. For breast cancer: 1) requested drug is being used for ovarian suppression in premenopausal patients and 2) the requested drug will be used in combination with endocrine therapy and 3) the disease is hormone receptor positive and 4) the disease is at a higher risk of recurrence (e.g., young age,

high-grade tumor, lymph-node involvement).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TREMFYA
Drug Names TREMFYA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For moderate to severe plaque psoriasis (new starts only): at least 3% of body surface

area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin,

intertriginous areas) are affected at the time of diagnosis.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupTRIKAFTADrug NamesTRIKAFTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For cystic fibrosis: The requested medication will not be used in combination with other

medications containing ivacaftor.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TRULICITY
Drug Names TRULICITY

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria The Prior Authorization only applies to patients whose claim is not submitted with an

ICD-10 code indicating a diagnosis of type 2 diabetes mellitus OR to patients who do not have a history of an antidiabetic drug (EXCLUDING glucagon-like peptide receptor agonists [GLP-1 RAs] and combination glucose-dependent insulinotropic polypeptide

[GIP] and GLP-1 RAs).

Prior Authorization Group TRUQAP
Drug Names TRUQAP

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group Drug Names PA Indication Indicator Off-label Uses TRUXIMA TRUXIMA

All FDA-approved Indications, Some Medically-accepted Indications Non-Hodgkin's lymphoma subtypes [small lymphocytic lymphoma (SLL), mantle cell lymphoma, marginal zone lymphomas (nodal, splenic, extranodal marginal zone lymphoma), Burkitt lymphoma, primary cutaneous B-cell lymphoma, high-grade B-cell lymphoma, histological transformation from indolent lymphomas to diffuse large B-cell lymphoma, histological transformation chronic lymphocytic leukemia (CLL)/SLL to diffuse large B-cell lymphoma. Castleman's disease, human immunodeficiency virus (HIV)-related B-cell lymphoma, hairy cell leukemia, post-transplant lymphoproliferative disorder (PTLD). B-cell lymphoblastic lymphomal, refractory immune or idiopathic thrombocytopenic purpura (ITP), autoimmune hemolytic anemia, Waldenstrom's macroglobulinemia/lymphoplasmacytic lymphoma, chronic graft-versus-host disease (GVHD), Sjogren syndrome, thrombotic thrombocytopenic purpura, refractory myasthenia gravis, Hodgkin's lymphoma (nodular lymphocyte-predominant), primary central nervous system (CNS) lymphoma, leptomeningeal metastases from lymphomas, acute lymphoblastic leukemia, prevention of Epstein-Barr virus (EBV)-related PTLD, multiple sclerosis, immune checkpoint inhibitor-related toxicities, pemphigus vulgaris, pediatric aggressive mature B-cell lymphomas, Rosai-Dorfman disease, and pediatric mature B-cell acute leukemia.

Exclusion Criteria
Required Medical Information

For moderately to severely active rheumatoid arthritis (new starts only): 1) patient meets ANY of the following: a) requested drug will be used in combination with methotrexate (MTX) OR b) patient has intolerance or contraindication to MTX, AND 2) patient meets ANY of the following: a) inadequate response, intolerance, or contraindication to MTX OR b) inadequate response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD.

Hematologic malignancies must be CD20-positive. For multiple sclerosis: 1) patient has a diagnosis of relapsing remitting multiple sclerosis, AND2) patient has had an inadequate response to two or more disease-modifying drugs indicated for multiple sclerosis despite adequate duration of treatment.

Age Restrictions
Prescriber Restrictions
Coverage Duration

Other Criteria

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Immune checkpoint inhibitor-related toxicities: 3 months, All other: Plan Year

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Prior Authorization Group TUKYSA Drug Names TUKYSA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent human epidermal growth factor receptor 2 (HER2)-positive breast cancer

Exclusion Criteria -

**Required Medical Information** For colorectal cancer (including appendiceal adenocarcinoma): 1) the patient has

advanced, unresectable, or metastatic disease AND 2) the patient has human epidermal growth factor receptor 2 (HER2)-positive disease AND 3) the patient has RAS wild-type disease AND 4) the requested drug will be used in combination with trastuzumab and 5) the patient has not previously been treated with a HER2 inhibitor.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group TURALIO
Drug Names TURALIO

**PA Indication Indicator**All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Langerhans cell histiocytosis, Erdheim-Chester disease, Rosai-Dorfman disease

Exclusion Criteria -

**Required Medical Information** For Langerhans cell histiocytosis: 1) disease has colony stimulating factor 1 receptor

(CSF1R) mutation. For Erdheim-Chester disease and Rosai-Dorfman disease: 1) disease has CSF1R mutation AND patient has any of the following: a) symptomatic

disease OR b) relapsed/refractory disease.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupUBRELVYDrug NamesUBRELVY

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For acute treatment of migraine: The patient has experienced an inadequate treatment

response, intolerance, or the patient has a contraindication to at least one triptan 5-HT1

receptor agonist.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** UPTRAVI

**Drug Names** UPTRAVI, UPTRAVI TITRATION PACK

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For pulmonary arterial hypertension (World Health Organization [WHO] Group 1): PAH

was confirmed by right heart catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment

pulmonary vascular resistance is greater than or equal to 3 Wood units.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** V-GO

**Drug Names** V-GO 20, V-GO 30, V-GO 40 **PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information

Omnipod GO, initial: 1) the patient has diabetes requiring insulin management AND 2) the patient is currently self-testing glucose levels, the patient will be counseled on self-testing glucose levels, or the patient is using a continuous glucose monitor AND 3) the patient has experienced an inadequate treatment response or intolerance to long-acting basal insulin therapy. Omnipod, V-GO, initial: 1) The patient has diabetes requiring insulin management with multiple daily injections AND 2) The patient is self-testing glucose levels 4 or more times per day OR the patient is using a continuous glucose monitor AND 3) The patient has experienced any of the following with the current diabetes regimen: inadequate glycemic control, recurrent hypoglycemia, wide fluctuations in blood glucose, dawn phenomenon with persistent severe early morning hyperglycemia, severe glycemic excursions.

Age Restrictions -

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VALCHLOR
Prug Names VALCHLOR

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Chronic or smoldering adult T-cell leukemia/lymphoma (ATLL), Stage 2 or higher

mycosis fungoides (MF)/Sezary syndrome (SS), primary cutaneous marginal zone lymphoma, primary cutaneous follicle center lymphoma, CD30-positive lymphomatoid papulosis (LvP), unifocal Langerhans cell histiocytosis (LCH) with isolated skin disease

Must have a documented diagnosis of a seizure disorder requiring acute treatment.

Exclusion Criteria -

Required Medical Information -

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VALTOCO

**Drug Names** VALTOCO 10 MG DOSE, VALTOCO 15 MG DOSE, VALTOCO 20 MG DOSE,

**VALTOCO 5 MG DOSE** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information

Required Medical Information

- - - -

Age Restrictions -

Prescriber Restrictions Must be prescribed by a neurologist or must be prescribed after consultation with a

neurologist.

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupVANFLYTADrug NamesVANFLYTA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group

**Drug Names** 

VENCLEXTA, VENCLEXTA STARTING PACK

PA Indication Indicator

All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Mantle cell lymphoma, blastic plasmacytoid dendritic cell neoplasm (BPDCN), multiple myeloma, relapsed or refractory acute myeloid leukemia (AML), Waldenstrom

myeloma, relapsed of refractory acute myeloid leukernia (AML), vvaluenstrom

macroglobulinemia/lymphoplasmacytic lymphoma, relapsed or refractory systemic light

chain amyloidosis with translocation t(11:14), myelodysplastic syndrome

**Exclusion Criteria** 

Required Medical Information

For acute myeloid leukemia (AML): 1) patient is 60 years of age or older, OR 2) patient is less than 60 years of age with unfavorable risk genetics and TP53-mutation, OR 3) patient has comorbidities that preclude use of intensive induction chemotherapy, OR 4) patient has relapsed or refractory disease. For blastic plasmacytoid dendritic cell

neoplasm (BPDCN): 1) patient has systemic disease being treated with palliative intent, OR 2) patient has relapsed or refractory disease. For multiple myeloma: 1) the disease is relapsed or progressive, AND 2) the requested drug will be used in combination with

dexamethasone, AND 3) patient has t(11:14) translocation. For Waldenstrom macroglobulinemia/lymphoplasmacytic lymphoma: 1) patient has previously treated

disease that did not respond to primary therapy, OR 2) patient has progressive or

relapsed disease.

**VENCLEXTA** 

Age Restrictions

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VENTAVIS
Drug Names VENTAVIS

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

Exclusion Criteria -

**Required Medical Information** For pulmonary arterial hypertension (World Health Organization [WHO] Group 1): PAH

was confirmed by right heart catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than 20 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment

pulmonary vascular resistance is greater than or equal to 3 Wood units.

**Age Restrictions** 

ge Restrictions -

Prescriber Restrictions

**Coverage Duration** 

Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization Group VERSACLOZ Drug Names VERSACLOZ

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the treatment of a severely ill patient with schizophrenia who failed to respond

adequately to standard antipsychotic treatment (i.e., treatment-resistant schizophrenia):
1) the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to one of the following generic products: aripiprazole, asenapine, lurasidone, olanzapine, quetiapine, risperidone, ziprasidone, AND 2) the patient has experienced an inadequate treatment response, intolerance, or has a contraindication

to one of the following brand products: Caplyta, Rexulti, Secuado, Vraylar.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VERZENIO
Drug Names VERZENIO

Drug Names VERZENIO
PA Indication Indicator All FDA-approx

PA Indication Indicator
All FDA-approved Indications, Some Medically-accepted Indications
Off-label Uses
Recurrent hormone receptor (HR)-positive, human epidermal growth factor receptor 2

(HER2)-negative breast cancer in combination with fulvestrant or an aromatase inhibitor, or as a single agent if progression on prior endocrine therapy and prior

chemotherapy in the metastatic setting.

Exclusion Criteria -

Required Medical Information

Age Restrictions

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupVICTOZADrug NamesVICTOZA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria - Required Medical Information - Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria The Prior Authorization only applies to patients whose claim is not submitted with an

ICD-10 code indicating a diagnosis of type 2 diabetes mellitus OR to patients who do not have a history of an antidiabetic drug (EXCLUDING glucagon-like peptide receptor agonists [GLP-1 RAs] and combination glucose-dependent insulinotropic polypeptide

[GIP] and GLP-1 RAs).

**Prior Authorization Group** VIGABATRIN

**Drug Names** VIGABATRIN, VIGADRONE, VIGPODER

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For complex partial seizures (i.e., focal impaired awareness seizures): patient has

experienced an inadequate treatment response to at least two antiepileptic drugs for

complex partial seizures (i.e., focal impaired awareness seizures).

Age Restrictions Infantile Spasms: 1 month to 2 years of age. Complex partial seizures (i.e., focal

impaired awareness seizures): 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupVIJOICEDrug NamesVIJOICE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -Required Medical Information -

**Age Restrictions** 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VITRAKVI
Drug Names VITRAKVI

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Non-metastatic neurotrophic tyrosine receptor kinase (NTRK) gene fusion-positive solid

tumors, first-line treatment of NTRK gene fusion-positive solid tumors.

Exclusion Criteria

**Required Medical Information** For all neurotrophic tyrosine receptor kinase (NTRK) gene fusion-positive solid tumors,

the disease is without a known acquired resistance mutation.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupVIZIMPRODrug NamesVIZIMPRO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent non-small cell lung cancer (NSCLC).

Exclusion Criteria -

**Required Medical Information** For non-small cell lung cancer (NSCLC): 1) the disease is recurrent, advanced or

metastatic, and 2) the patient has sensitizing EGFR mutation-positive disease.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VONJO
Drug Names VONJO

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupVORICONAZOLEDrug NamesVORICONAZOLE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** The patient will use the requested drug orally or intravenously.

Age Restrictions Prescriber Restrictions -

Coverage Duration 6 months

Other Criteria -

Prior Authorization GroupVOSEVIDrug NamesVOSEVI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

**Exclusion Criteria** Decompensated cirrhosis/moderate or severe hepatic impairment (Child Turcotte Pugh

class B or C)

**Required Medical Information** For hepatitis C: Infection confirmed by presence of HCV RNA in the serum prior to

starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated substitutions where applicable, transplantation status if applicable. Coverage conditions and specific durations of approval will be based on

current American Association for the Study of Liver Diseases and Infectious Diseases

Society of America (AASLD-IDSA) treatment guidelines.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Criteria will be applied consistent with current AASLD-IDSA guidance.

Other Criteria -

**Prior Authorization Group** VOTRIENT

**Drug Names** PAZOPANIB HYDROCHLORIDE, VOTRIENT

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Thyroid carcinoma (follicular, papillary, Hurthle cell, or medullary), uterine sarcoma,

chondrosarcoma, gastrointestinal stromal tumor

Exclusion Criteria

**Required Medical Information** For renal cell carcinoma: 1) The disease is advanced, relapsed, or stage IV, OR 2) the

requested drug will be used for von Hippel-Lindau (VHL)-associated renal cell carcinoma. For gastrointestinal stromal tumor (GIST): the patients meets one of the following: 1) the disease is unresectable, recurrent/progressive, or metastatic AND the patient has failed an FDA-approved therapy (e.g., imatinib, sunitinib, regorafenib, ripretinib), 2) the requested drug will be used for unresectable succinate

dehydrogenase (SDH)-deficient GIST, OR 3) the requested drug will be used for the palliation of symptoms if previously tolerated and effective. For soft tissue sarcoma

(STS): The patient does not have an adipocytic soft tissue sarcoma. For uterine

sarcoma: The disease is recurrent or metastatic.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group VOWST

Drug Names VOWST

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the prevention of recurrence of Clostridioides difficile infection (CDI): 1) The

diagnosis of CDI has been confirmed by a positive stool test for C. difficile toxin, AND 2) The requested drug will be administered at least 48 hours after the last dose of

antibiotics used for the treatment of recurrent CDI.

**Age Restrictions** 18 years of age or older

Prescriber Restrictions -

Coverage Duration 1 month

Other Criteria -

Prior Authorization Group VYNDAMAX Drug Names VYNDAMAX

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

Required Medical Information F

For cardiomyopathy of hereditary or wild-type transthyretin-mediated amyloidosis (ATTR-CM): Initiation: 1) patient exhibits clinical manifestation of disease (e.g., dyspnea, fatigue, orthostatic hypotension, syncope, peripheral edema), AND 2) cardiac involvement was confirmed by echocardiography or cardiac magnetic resonance imaging (e.g., end-diastolic interventricular septal wall thickness exceeding 12 millimeters), AND 3) patient meets one of the following: a) if the request is for hereditary ATTR-CM the patient is positive for a mutation of the transthyretin (TTR) gene, b) if the request is for wild-type ATTR-CM the patient has transthyretin precursor proteins confirmed by testing. Continuation: patient demonstrates a beneficial response to therapy (e.g., slowing of clinical decline).

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupVYNDAQELDrug NamesVYNDAQEL

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** 

For cardiomyopathy of hereditary or wild-type transthyretin-mediated amyloidosis (ATTR-CM): Initiation: 1) patient exhibits clinical manifestation of disease (e.g., dyspnea, fatigue, orthostatic hypotension, syncope, peripheral edema), AND 2) cardiac involvement was confirmed by echocardiography or cardiac magnetic resonance imaging (e.g., end-diastolic interventricular septal wall thickness exceeding 12 millimeters), AND 3) patient meets one of the following: a) if the request is for hereditary ATTR-CM the patient is positive for a mutation of the transthyretin (TTR) gene, b) if the request is for wild-type ATTR-CM the patient has transthyretin precursor proteins confirmed by testing. Continuation: patient demonstrates a beneficial response to therapy (e.g., slowing of clinical decline).

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group WAINUA Drug Names** WAINUA

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** For polyneuropathy of hereditary transthyretin (TTR)-mediated amyloidosis, initial

> therapy: Patient is positive for a mutation of the TTR gene and exhibits clinical manifestation of disease (for example, amyloid deposition in biopsy specimens, TTR protein variants in serum, progressive peripheral sensory-motor polyneuropathy). For polyneuropathy of hereditary TTR-mediated amyloidosis, continuation: Patient demonstrates a beneficial response to therapy (for example, improvement of

neuropathy severity and rate of disease progression).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** Plan Year

Other Criteria

**Prior Authorization Group WELIREG WELIREG Drug Names** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria** 

**Required Medical Information** For advanced renal cell carcinoma (RCC): 1) patient previously received treatment with

> a programmed death receptor-1 (PD-1) or programmed death-ligand 1 (PD-L1) inhibitor, AND 2) patient previously received treatment with a vascular endothelial growth factor tyrosine kinase inhibitor (VEGF-TKI) [e.g., Cabometyx (cabozantinib),

Inlyta (axitinib), Nexavar (sorafenib)].

Age Restrictions

**Prescriber Restrictions** Plan Year

**Coverage Duration** 

Other Criteria

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Prior Authorization Group WINLEVI
Drug Names WINLEVI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** The patient has experienced an inadequate treatment response, intolerance or the

patient has a contraindication to a generic acne product (e.g., topical clindamycin,

topical erythromycin, topical retinoid, or oral isotretinoin).

**Age Restrictions** 12 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group XALKORI Drug Names XALKORI

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent non-small cell lung cancer (NSCLC), NSCLC with high-level MET

amplification or MET exon 14 skipping mutation, symptomatic or relapsed/refractory anaplastic lymphoma kinase (ALK)-fusion positive Erdheim-Chester Disease, symptomatic or relapsed/refractory (ALK)-fusion positive Rosai-Dorfman Disease,

(ALK)-fusion positive Langerhans Cell Histiocytosis.

Exclusion Criteria -

**Required Medical Information** For NSCLC, the requested drug is used in any of the following settings: 1) the patient

has recurrent, advanced or metastatic ALK-positive NSCLC, OR 2) the patient has recurrent, advanced or metastatic ROS-1 positive NSCLC, OR 3) the patient has NSCLC with high-level MET amplification or MET exon 14 skipping mutation. For IMT,

the disease is ALK-positive. For ALCL, the disease is relapsed or refractory and

ALK-positive.

Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** XELJANZ

Drug NamesXELJANZ, XELJANZ XRPA Indication IndicatorAll FDA-approved Indications

Off-label Uses - Exclusion Criteria -

Required Medical Information

For moderately to severely active rheumatoid arthritis (new starts only): patient has experienced an inadequate treatment response, intolerance or has a contraindication to at least one tumor necrosis factor (TNF) inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]), For active psoriatic arthritis (new starts only): 1) patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one TNF inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]) AND 2) the requested drug is used in combination with a nonbiologic DMARD. For active ankylosing spondylitis (new starts only): Patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one tumor necrosis factor (TNF) inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]). For moderately to severely active ulcerative colitis (new starts only): patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one tumor necrosis factor (TNF) inhibitor (e.g., adalimumab-aacf, Humira [adalimumab], Idacio [adalimumab-aacf]). For active polyarticular course juvenile idiopathic arthritis (new starts only): patient has experienced an inadequate treatment response, intolerance, or has a contraindication to at least one tumor necrosis factor (TNF) inhibitor (e.g., adalimumab-aacf, Enbrel [etanercept], Humira [adalimumab], Idacio [adalimumab-aacf]).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupXERMELODrug NamesXERMELO

PA Indication Indicator All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** XGEVA **Drug Names** XGEVA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** For hypercalcemia of malignancy: condition is refractory to intravenous (IV)

bisphosphonate therapy or there is a clinical reason to avoid IV bisphosphonate

therapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization Group XHANCE Drug Names XHANCE

PA Indication Indicator All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Patient has experienced an inadequate treatment response to generic fluticasone nasal

spray.

Age Restrictions 18 years of age or older

Prescriber Restrictions

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group XIFAXAN Drug Names XIFAXAN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For irritable bowel syndrome with diarrhea (IBS-D): 1) The patient has not previously

received treatment with the requested drug OR 2) The patient has previously received treatment with the requested drug AND a) the patient is experiencing a recurrence of symptoms AND b) the patient has not already received an initial 14-day course of treatment and two additional 14-day courses of treatment with the requested drug.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Reduction in risk of overt HE recurrence: 6 Months, IBS-D: 14 Days

Other Criteria -

**Prior Authorization Group** XIFAXAN 200MG

**Drug Names** XIFAXAN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of Traveler's diarrhea caused by Noninvasive

strains of E coli.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group XOLAIR XOLAIR Drug Names** 

PA Indication Indicator All FDA-approved Indications

Off-label Uses **Exclusion Criteria Required Medical Information** 

For moderate to severe persistent asthma, initial therapy: 1) Patient has a positive skin test (or blood test) to at least one perennial aeroallergen, 2) Patient has baseline IgE level greater than or equal to 30 IU/mL, AND 3) Patient has inadequate asthma control despite current treatment with both of the following medications: a) Medium-to-high-dose inhaled corticosteroid, AND b) Additional controller (i.e., long acting beta2-agonist, long-acting muscarinic antagonist, leukotriene modifier, or sustained-release theophylline) unless patient has an intolerance or contraindication to such therapies. For moderate to severe persistent asthma, continuation of therapy: Asthma control has improved on treatment with the requested drug, as demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations or a reduction in the daily maintenance oral corticosteroid dose. For chronic spontaneous urticaria (CSU), initial therapy: 1) Patient has been evaluated for other causes of urticaria, including bradykinin-related angioedema and IL-1-associated urticarial syndromes (e.g., auto-inflammatory disorders, urticarial vasculitis), 2) Patient has experienced a spontaneous onset of wheals, angioedema, or both, for at least 6 weeks, AND 3) Patient remains symptomatic despite H1 antihistamine treatment. For CSU. continuation of therapy: Patient has experienced a benefit (e.g., improved symptoms) since initiation of therapy. For chronic rhinosinusitis with nasal polyps (CRSwNP): 1) The requested drug is used as add-on maintenance treatment, AND 2) Patient has experienced inadequate treatment response to Xhance (fluticasone). CSU: 12 years of age or older. Asthma: 6 years of age or older. CRSwNP: 18 years of

Age Restrictions

**Prescriber Restrictions Coverage Duration** Other Criteria

age or older. IgE-mediated food allergy: 1 year of age or older

CSU initial: 6 months, All others: Plan Year

For IgE-mediated food allergy, initial therapy: Patient has baseline IgE level greater than or equal to 30 IU/mL. For IgE-mediated food allergy, continuation of therapy: Patient has experienced a benefit as evidenced by a decrease in hypersensitivity (e.g., moderate to severe skin, respiratory or gastrointestinal symptoms) to food allergen.

Updated 07/01/2024 147 Prior Authorization Group XOSPATA
Drug Names XOSPATA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia and FLT3

rearrangement

Exclusion Criteria

**Required Medical Information** For myeloid, lymphoid, or mixed lineage neoplasms with eosinophilia and FMS-like

tyrosine kinase 3 (FLT3) rearrangement: the disease is in chronic or blast phase.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** XPOVIO

Drug Names XPOVIO, XPOVIO 60 MG TWICE WEEKLY, XPOVIO 80 MG TWICE WEEKLY

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses

Histologic transformation of indolent lymphomas to diffuse large B-cell lymphoma, acquired immunodeficiency syndrome (AIDS)-related B-cell lymphoma, high-grade

B-cell lymphoma

Exclusion Criteria -

**Required Medical Information** For multiple myeloma: Patient must have been treated with at least one prior therapy.

For B-cell lymphomas: Patient must have been treated with at least two lines of

systemic therapy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group XTANDI Drug Names XTANDI

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -

Exclusion Criteria -

**Required Medical Information** For the treatment of castration-resistant prostate cancer or metastatic

castration-sensitive prostate cancer: The requested drug will be used in combination with a gonadotropin-releasing hormone (GnRH) analog or after bilateral orchiectomy.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group XURIDEN Drug Names XURIDEN

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** Must have a documented diagnosis of hereditary orotic aciduria.

Age Restrictions - Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

**Prior Authorization Group** XYREM

**Drug Names** SODIUM OXYBATE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** 

For the treatment of excessive daytime sleepiness in a patient with narcolepsy, initial request: 1) The diagnosis has been confirmed by sleep lab evaluation, AND 2) The patient meets one of the following criteria: a) if the patient is 17 years of age or younger, the patient has experienced an inadequate treatment response or intolerance to at least one central nervous system (CNS) stimulant drug (e.g., amphetamine. dextroamphetamine, methylphenidate). OR has a contraindication that would prohibit a trial of central nervous system (CNS) stimulant drugs (e.g., amphetamine, dextroamphetamine, methylphenidate), b) If the patient is 18 years of age or older, the patient has experienced an inadequate treatment response or intolerance to at least one central nervous system (CNS) wakefulness promoting drug (e.g., armodafinil, modafinil), OR has a contraindication that would prohibit a trial of central nervous system (CNS) wakefulness promoting drugs (e.g., armodafinil, modafinil). For the treatment of cataplexy in a patient with narcolepsy, initial request: The diagnosis has been confirmed by sleep lab evaluation. If the request is for a continuation of therapy. then the patient experienced a decrease in daytime sleepiness with narcolepsy or a decrease in cataplexy episodes with narcolepsy.

Age Restrictions 7 years of age or older

Prescriber Restrictions Prescribed by or in consultation with a sleep disorder specialist or neurologist

Coverage Duration Plan Year

Other Criteria

Prior Authorization Group XYWAV
Drug Names XYWAV

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses
Exclusion Criteria
Required Medical Information

For the treatment of excessive daytime sleepiness in a patient (pt) with narcolepsy, initial request: 1) the diagnosis (dx) has been confirmed by sleep lab evaluation, AND 2) the pt meets one of the following criteria: a) If the pt is 17 years of age or younger. the pt has experienced an inadequate treatment response or intolerance to at least one central nervous system (CNS) stimulant drug (e.g., amphetamine, dextroamphetamine, methylphenidate). OR has a contraindication that would prohibit a trial of CNS stimulant drugs (e.g., amphetamine, dextroamphetamine, methylphenidate), b) If the pt is 18 years of age or older, the pt has experienced an inadequate treatment response or intolerance to at least one CNS wakefulness promoting drug (e.g., armodafinil, modafinil). OR has a contraindication that would prohibit a trial of CNS wakefulness promoting drugs (e.g., armodafinil, modafinil). For idiopathic hypersomnia the diagnosis has been confirmed by ALL of the following: 1) pt has experienced lapses into sleep or an irrepressible need to sleep during daytime, on a daily basis, for at least 3 months. AND 2) insufficient sleep syndrome is confirmed absent, AND 3) cataplexy is absent, AND 4) fewer than 2 sleep onset rapid eye movement periods (SOREMPs) or no SOREMPs, if the rapid eye movement latency on an overnight sleep study was less than or equal to 15 minutes, AND 5) average sleep latency of less than or equal to 8 minutes on Multiple Sleep Latency Test or total 24-hour sleep time is greater than or equal to 11 hours, AND 6) another condition (sleep disorder, medical or psychiatric disorder, or drug/medication use) does not better explain the hypersomnolence and test results.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Narcolepsy: 7 years of age or older, Idiopathic hypersomnia: 18 years of age or older Prescribed by or in consultation with a sleep disorder specialist or neurologist Plan Year

For the treatment of cataplexy in a pt with narcolepsy, initial request: the dx has been confirmed by sleep lab evaluation. For narcolepsy, continuation of therapy: the pt has experienced a decrease in daytime sleepiness with narcolepsy or a decrease in cataplexy episodes with narcolepsy. For idiopathic hypersomnia, continuation of therapy: the pt has experienced a decrease in daytime sleepiness from baseline.

Prior Authorization Group YONSA
Drug Names YONSA

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -

**Required Medical Information** The requested drug will be used in combination with a gonadotropin-releasing hormone

(GnRH) analog or after bilateral orchiectomy.

Age Restrictions -Prescriber Restrictions --

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupZARXIODrug NamesZARXIO

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Neutropenia in myelodysplastic syndromes (MDS), agranulocytosis, neutropenia in

aplastic anemia, human immunodeficiency virus (HIV)-related neutropenia, neutropenia

related to renal transplant, hematopoietic syndrome of acute radiation syndrome

**Exclusion Criteria**Use of the requested product within 24 hours prior to or following chemotherapy. **Required Medical Information**For prophylaxis or treatment of myelosuppressive chemotherapy-induced febrile

neutropenia (FN) patient must meet both of the following: 1) Patient has a solid tumor or non-myeloid cancer, and 2) Patient has received, is currently receiving, or will be

receiving treatment with myelosuppressive anti-cancer therapy.

Age Restrictions - Prescriber Restrictions -

Coverage Duration 6 months

Other Criteria -

Prior Authorization GroupZEJULADrug NamesZEJULA

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Uterine leiomyosarcoma

Exclusion Criteria -

**Required Medical Information** For uterine leiomyosarcoma: 1) the requested drug is used as second-line therapy AND

2) the patient has BRCA-altered disease.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ZELBORAF
Drug Names ZELBORAF

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Non-small cell lung cancer, hairy cell leukemia, thyroid carcinoma (i.e., papillary

carcinoma, follicular carcinoma, and Hurthle cell carcinoma), central nervous system cancer (i.e., glioma, astrocytoma, glioblastoma, pediatric diffuse high-grade glioma),

adjuvant systemic therapy for cutaneous melanoma, Langerhans cell histiocytosis.

Exclusion Criteria

Required Medical Information For central nervous system (CNS) cancer (i.e., glioma, astrocytoma, glioblastoma,

pediatric diffuse high-grade glioma): 1) The tumor is positive for BRAF V600E mutation, AND 2) The requested drug will be used in combination with cobimetinib OR the requested drug is being used for the treatment of pediatric diffuse high-grade glioma. For melanoma: 1) The tumor is positive for BRAF V600 activating mutation (e.g., V600E or V600K), AND 2) the requested drug will be used as a single agent, or in combination with cobimetinib, AND 3) The requested drug will be used for either of the following: a) unresectable, limited resectable, or metastatic disease, or b) adjuvant systemic therapy. For Erdheim-Chester Disease and Langerhans Cell Histiocytosis: Tumor is positive for BRAF V600 mutation. For non-small cell lung cancer: 1) The tumor is positive for the BRAF V600E mutation, AND 2) The patient has recurrent, advanced, or metastatic disease. For papillary, follicular, and hurthle cell thyroid carcinoma: 1) The tumor is positive for BRAF mutation, AND 2) The disease is not

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ZIEXTENZO

Drug Names ZIEXTENZO

PA Indication Indicator All FDA-approved Indication

PA Indication Indicator All FDA-approved Indications, Some Medically-accepted Indications
Off-label Uses Stem cell transplantation-related indications

Exclusion Criteria Use of the requested product less than 24 hours before or after chemotherapy.

amenable to radioactive iodine (RAI) therapy.

**Required Medical Information**For prophylaxis of myelosuppressive chemotherapy-induced febrile neutropenia: the patient must meet both of the following: 1) Patient has a solid tumor or non-myeloid cancer, and 2) Patient is currently receiving or will be receiving treatment with

myelosuppressive anti-cancer therapy.

Age Restrictions -

**Prescriber Restrictions** 

Coverage Duration 6 months

Other Criteria -

Prior Authorization GroupZILBRYSQDrug NamesZILBRYSQ

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses Exclusion Criteria Required Medical Information Age Restrictions Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Off-label Uses

Prior Authorization GroupZIRABEVDrug NamesZIRABEV

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Ampullary adenocarcinoma, breast cancer, central nervous system (CNS) cancers, malignant pleural mesothelioma, malignant peritoneal mesothelioma, pericardial mesothelioma, tunica vaginalis testis mesothelioma, soft tissue sarcomas, uterine neoplasms, endometrial carcinoma, vulvar cancers, small bowel adenocarcinoma, and ophthalmic-related disorders: diabetic macular edema, neovascular (wet) age-related macular degeneration including polypoidal choroidopathy and retinal angiomatous proliferation subtypes, macular edema following retinal vein occlusion, proliferative diabetic retinopathy, choroidal neovascularization, neovascular glaucoma and

retinopathy of prematurity.

Exclusion Criteria -

Required Medical Information -

Age Restrictions -

Prescriber Restrictions - Coverage Duration Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization GroupZOLINZADrug NamesZOLINZA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Mycosis fungoides (MF)/Sezary syndrome (SS)

Exclusion Criteria -

Required Medical Information -

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupZONISADEDrug NamesZONISADE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For adjunctive treatment of partial-onset seizures (i.e., focal-onset seizures): 1) The

patient has experienced an inadequate treatment response, intolerance, or has a contraindication to a generic anticonvulsant AND the patient has experienced an inadequate treatment response, intolerance, or has a contraindication to any of the following: Aptiom, Xcopri, Spritam OR 2) The patient has difficulty swallowing solid oral

dosage forms (e.g., tablets, capsules).

**Age Restrictions** 16 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ZTALMY
Drug Names ZTALMY

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses -Exclusion Criteria -Required Medical Information -

**Age Restrictions** 2 years of age or older

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupZURZUVAEDrug NamesZURZUVAE

**PA Indication Indicator** All FDA-approved Indications

Off-label Uses - Exclusion Criteria -

**Required Medical Information** For the treatment of postpartum depression (PPD): diagnosis was confirmed using

standardized rating scales that reliably measure depressive symptoms (e.g., Hamilton Depression Rating Scale [HDRS], Edinburgh Postnatal Depression Scale [EPDS], Patient Health Questionnaire 9 [PHQ9], Montgomery-Asberg Depression Rating Scale

[MADRS], Beck's Depression Inventory [BDI], etc.).

Age Restrictions -

Prescriber Restrictions -

Coverage Duration 1 month

Other Criteria -

Prior Authorization GroupZYDELIGDrug NamesZYDELIG

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Small lymphocytic lymphoma (SLL)

Exclusion Criteria -

**Required Medical Information** For CLL/SLL: the requested drug is used as second-line or subsequent therapy

Age Restrictions

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization Group ZYKADIA
Prug Names ZYKADIA

**PA Indication Indicator** All FDA-approved Indications, Some Medically-accepted Indications

Off-label Uses Recurrent ALK-positive non-small cell lung cancer (NSCLC), recurrent, advanced, or

metastatic ROS1-positive NSCLC, inflammatory myofibroblastic tumor (IMT), brain

metastases from NSCLC.

Exclusion Criteria

**Required Medical Information** For NSCLC: the patient has recurrent, advanced, or metastatic ALK-positive or

ROS1-positive disease. For inflammatory myofibroblastic tumor: the disease is

ALK-positive. For brain metastases from NSCLC: the patient has ALK-positive NSCLC.

Age Restrictions -

Prescriber Restrictions -

Coverage Duration Plan Year

Other Criteria -

Prior Authorization GroupZYPREXA RELPREVVDrug NamesZYPREXA RELPREVV

PA Indication Indicator All FDA-approved Indications

Off-label Uses -

Exclusion Criteria -

**Required Medical Information** Tolerability with oral olanzapine has been established.

Coverage Duration Plan Year

Other Criteria -